

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

- (51) International Patent Classification 5: C07C 323/62, 323/60, C07D 333/34 C07C 327/32, 317/50, 313/48 A61K 31/13, 31/38
- (11) International Publication Number:

WO 90/05719

A1 (43) International Publication Date: . 31 May 1990 (31.05.90)

(21) International Application Number:

PCT/GB89/01399

(22) International Filing Date:

23 November 1989 (23.11.89)

(30) Priority data:

8827305.7

23 November 1988 (23.11.88) GB

- (71) Applicant (for all designated States except US): BRITISH BIO-TECHNOLOGY LIMITED [GB/GB]; Watlington Road, Cowley, Oxford OX4 5LY (GB).
- (72) Inventors; and
- (72) Inventors; and
 (75) Inventors/Applicants (for US only): CAMPION, Colin [GB/GB]; 3 Howe Close, Wheatley, Oxon OX4 5LY (GB). DAVIDSON, Alan, Hornsby [GB/GB]; 27 Newland Mill, Witney, Oxon OX8 6HH (GB). DICKENS, Jonathan, Philip [GB/GB]; Burton House, Park Farm Road, High Wycombe, Bucks HP12 4AF (GB). CRIMMIN, Michael, John [GB/GB]; Oaklea, 64 Fernbank Road, Ascot SL5 8HE (GB).

- (74) Agents: SHEARD, Andrew, Gregory et al.; Kilburn & Strode, 30 John Street, London WC1N 2DD (GB).
- (81) Designated States: AT (European patent), AU, BE (European patent), CH (European patent), DE (European patent), DK, ES (European patent), FI, FR (European patent), GB (European patent), IT (European patent), JP, LU (European patent), NL (European patent), NO, SE (European patent), US.

Published

With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: HYDROXAMIC ACID BASED COLLAGENASE INHIBITORS

(57) Abstract

Compounds of general formula (I), wherein R1 represents hydrogen or an alkyl, phenyl, thiophenyl, substituted phenyl, phenylalkyl, heterocyclyl, alkylcarbonyl phenacyl or substituted phenacyl group; or, when n = 0, R^1 represents SR^X , wherein RX represents a group (α); R2 represents a hydrogen atom or an alkyl, alkenyl, phenylalkyl, cycloalkylalkyl or cycloalkenylalkyl group; R3 represents an amino acid residue with R or S stereochemistry or an alkyl, benzyl, (C1-C6 alkoxy) benzyl or benzyloxy(C₁-C₆ alkyl) group; R⁴ represents a hydrogen atom or an alkyl group; R⁵ represents a hydrogen atom or a methyl group; n is an integer having the value 0, 1 or 2; and A represents a hydrocarbon chain optionally substituted with one or more alkyl, phenyl or substituted phenyl groups; and their salts and N-oxides are collagenase inhibitors and are useful in the management of disease involving tissue degradation and/or the promotion of wound healing. Diseases involving tissue degradation include arthropathy (particularly rheumatoid arthritis), inflammation, dermatological diseases, bone resorption diseases and tumour invasion.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

		•				
	AT	Austria	ES:	Spain	MG	Madagascar
	AU	Australia	FT	Finland	ML	Mali
	BB	Barbados	FR	France	MR	Mauritania
	BE	Belgium	GA	Gabon:	MW	Malawi
	BF	Burkina Fasso	GB	United Kingdom	NL	Netherlands
•	BG	Bulgaria	HU	Hungary	NO	Norway
	BJ	Benin	IT	Italy	. RO	Romania
	BR -	Brazil	JР	Japan	ŚD	Sudan
	CA	Canada	KP	Democratic People's Republic	SE	Sweden
	CF	Central African Republic		of Korea	SN	Senegal
	CG	Congo	- KR	Republic of Korea	SU	Soviet Union
	СН	Switzerland	П	Liechtenstein	TD	Chad
	CM	Cameroon	IK	Sri Lanka	TG	Togo
	DE	Germany, Federal Republic of	ī	Luxembourg .	US	United States of Am
	ناب	Carmeil's anima scalange or				

1

1 HYDROXAMIC ACID BASED COLLAGENASE INHIBITORS.

2 .

This invention relates to pharmaceutically and veterinarily active compounds, which are derivatives of hydroxamic acid.

6

The compounds of the present invention act as 7 inhibitors of metalloproteases involved in tissue 8 degradation, such as collagenase, which initiates 9 collagen breakdown, stromelysin (protoglycanase), 10 gelatinase and collagenase (IV). There is evidence 11 implicating collagenase as one of the key enzymes in 12 articular cartilage and bone in breakdown of 13 rheumatoid arthritis (Arthritis and Rheumatism, 20, 14 1231 - 1239, 1977). Potent inhibitors of collagenase 15 and other metalloproteases involved in tissue 16 degradation are useful in the treatment of rheumatoid 17 arthritis and related diseases in which collagenolytic 18 activity is important. Inhibitors of metalloproteases 19 of this type can therefore be used in treating or 20 preventing conditions which involve tissue breakdown; 21 they are therefore useful in the treatment of 22 arthropathy, dermatological conditions, bone 23 resorption, inflammatory diseases and tumour invasion 24 and in the promotion of wound healing. Specifically, 25 compounds of the present invention may be useful in the 26 treatment of osteopenias such as osteoporosis, 27 rheumatoid arthritis, osteoarthritis, periodontitis, 28 gingivitis, corneal ulceration and tumour invasion. 29

30

A number of small peptide like compounds which inhibit metalloproteases have been described. Perhaps the most notable of these are those relating to the WO 90/05719 .

31 32 33 2

angiotensin converting enzyme (ACE) where 1 such 2 agents act to block the conversion of the decapeptide angiotensin I to angiotensin II a potent pressor 3 substance. Compounds of this type are described in 4 EP-A-0012401: 5 6 7 hydroxamic acids have been suggested as Certain 8 collagenase inhibitors as in US-A-4599361 and 9 EP-A-0236872. Other hydroxamic acids have been prepared 10 as ACE inhibitors, for example in US-A-4105789, while still others have been described 11 as enkephalinase inhibitors as in US-A-4496540. 12 13 14 EP-A-0012401 discloses antihypertensive compounds of 15 the formula: 16 OR^1 R^3 R^4 R^5 O17 18 " | 1" R-C-C-NH-CH-C-N--C-R 19 20 \mathbb{R}^2 R^7 21. 0 22 23 wherein 24 R and R⁶ are the same or different and are hydroxy, 25 26 alkoxy, alkenoxy, dialkylamino alkoxy, acylamino alkoxy, acyloxy alkoxy, aryloxy, alkyloxy, substituted 27 aryloxy or substituted aralkoxy wherein the substituent 28 is methyl, halo, or methoxy, amino, alkylamino, 29 30 dialkylamino, aralkylamino or hydroxyamino;

```
R<sup>1</sup> is hydrogen, alkyl of from 1 to 20 carbon atoms,
    including branched, cyclic and unsaturated alkyl
    groups;
 3
 4
    substituted alkyl wherein the substituent is halo,
5
    hydroxy, alkoxy, aryloxy amino, alkylamino,
6
    dialkylamino, acrylamino, arylamino, guanidino,
7
    imidazolyl, indolyl, mercapto, alkylthio, arylthio,
8
    carboxy, carboxamido, carbalkoxy, phenyl, substituted
9
    phenyl wherein the substituent is alkyl, alkoxy or
10
    halo; aralkyl or heteroaralkyl, aralkenyl or
11
    heteroaralkenyl, substituted aralkyl, substituted
12
    heteroaralkyl, substituted aralkenyl or substituted
13
    hetereoaralkenyl, wherein the substituent is halor or
14
    dihalo, alkyl, hydroxy, alkoxy, amino, aminomethyl,
15
    acrylamino, dialkylamino, alkylamino, carboxyl,
16
    haloalkyl, cyano or sulphonamido, aralkyl or
17
    hetereoaralkyl substituted on the alkyl portion by
18
    amino or acylamino;
19
20
    R<sup>2</sup> and R<sup>7</sup> are hydrogen or alkyl;
21
22
         is hydrogen, alkyl, phenylalkyl,
23
    \mathbf{R}^3
    aminomethylphenylalkyl, hydroxyphenylalkyl,
24
    hydroxyalkyl, acetylaminoalkyl, acylaminoalkyl,
25
    acylaminoalkyl aminoalkyl, dimethylaminoalkyl,
26
    haloalkyl, guanidinoalkyl, imidazolylalkyl,
27
    indolylalkyl, mercaptoalkyl and alkylthioalkyl;
28
29
    R4 is hydrogen or alkyl;
30
31
32
33
```

WO 90/05719

```
is hydrogen,
                         alkyl, phenyl, phenylalkyl,
2
    hydroxyphenylalkyl, hydroxyalkyl, aminoalkyl,
    guanidinoalkyl, imidazolylalkyl, indolylalkyl,
    mercaptoalkyl or alkylthioalkyl;
5
    R4 and R5 may be connected together to form an alkylene
6
    bridge of from 2 to 4 carbon atoms, an alkylene bridge
    of from 2 to 3 carbon atoms and one sulphur atom, an
8
    alkylene bridge of from 3 to 4 carbon atoms containing
9
    a double bond or an alkylene bridge as above,
10
    substituted with hydroxy, alkoxy or alkyl and the
11
12
    pharmaceutically acceptable salts thereof.
13
14
    US-A-4599361 discloses compounds of the formula:
15
16
                    HOHNC-A-CNH-CH-CNHR<sup>1</sup>
17
18
19
    wherein
20
    R^1 is C_1 - C_6 alkyl;
21
    R^2 is C_1-C_6 alkyl, benzyl, benzyloxybenzyl, (C_1-C_6)
22
     alkoxy)benzyl or benzyloxy(C1-C6 alkyl);
23
     a is a chiral centre with optional R or S
24
     stereochemistry;
25
     A is a
26
                    -(CHR^3-CHR^4)-group
27
28
29
     or a -(CR^3=CR^4) - group wherein b and c are chiral
30
     centres with optional R or S stereochemistry;
31
32
33
```

 R^3 is hydrogen, C_1 - C_6 alkyl, phenyl or phenyl(C_1 - C_6 alkyl) and R^4 is hydrogen, C_1 - C_6 alkyl, phenyl(C_1 - C_6 alkyl), cycloalkyl or cycloalkyl(C_1 - C_6 alkyl).

EP-A-0236872 discloses generically compounds of the formula

13 wherein

A represents a group of the formula HN(OH)-CO- or HCO-N(OH)-;

 R^1 represents a C_2 - C_5 alkyl group;

R² represents the characterising group of a natural alpha-amino acid in which the functional group can be protected, amino groups may be acylated and carboxyl groups can be amidated, with the proviso that R² can not represent hydrogen or a methyl group;

 R^3 represents hydrogen or an amino, hydroxy, mercapto, c_1-c_6 alkyl, c_1-c_6 alkoxy, c_1-c_6 acylamino, $c_1-c_6-alkyl$ thio, aryl- $(c_1-c_6-alkyl)-$, amino- $(c_1-c_6-alkyl)-$, hydroxy $(c_1-c_6-alkyl)-$, mercapto $(c_1-c_6-alkyl)$ or carboxy $(c_1-c_6-alkyl)$ group,

wherein the amino, hydroxy, mercapto or carboxyl groups 1 2 . can be protected and the amino groups may be acylated or the carboxyl groups may be amidated; 3 4 R4 represents hydrogen or a methyl group; 6 7 R^5 represents hydrogen or a C_1-C_6 acyl, C_1-C_6 alkoxy c_1-c_6 alkyl, $di(c_1-c_6-alkoxy)$ methylene, carboxy, (c_1-c_6) 8 alkyl)carbinyl, (C1-C6 alkoxy)carbinyl, arylmethoxy 9 carbinyl, (C1-C6 alkyl)amino carbinyl or arylamino 10 carbinyl group; and 11 12 R⁶ represents hydroxy or a methylene group; or 13 14 \mathbb{R}^2 and \mathbb{R}^4 together represent a group-(CH₂)_n-, wherein n 15 represents a number from 4 to 11; or 16 17 R4 and R5 together represent a trimethylene group; 18 19 20 and pharmaceutically acceptable salts of such 21 compounds, which are acid or basic. 22 US-A-4105789 generically discloses compounds which have 23 the general formula 24 25 26 R_4 -oc-(CH₂)_n-CH-CO-N-CH-COOH 27 28 and salts thereof, wherein 29 30 is hydrogen, lower alkyl, phenyl lower alkylene, 31 hydroxy-lower alkylene, hydroxyphenyl lower 32 alkylene, amino-lower alkylene, guanidine lower 33 .

7

alkylene, mercapto-lower alkylene, 1 alkyl-mercapto-lower alkylene, imidazolyl lower indolyl-lower alkylene or carbamoyl alkylene, 3 lower alkylene; 4 is hydrogen or lower alkyl; 5 R_2 is lower alkyl or phenyl lower alkylene; 6 R_2 is hydroxy, lower alkoxy or hydroxyamino; and 7 R_{Δ} 8 is 1 or 2. n 9 US-A-4496540 discloses compounds of the general 10 formula: 11 12 13 A-B-NHOH 14 wherein A is one of the aromatic group-containing amino 15 acid residues L-tryptophyl, D-tryptophyl, L-tyrosyl, 16 D-tyrosyl, L-phenylalanyl, or D-phenylalanyl, and B is 17 one of the amino acids glycine, L-alanine, D-alanine, 18 L-leucine, D-leucine, L-isoleucine, or D-isoleucine; 19 and pharmaceutically acceptable salts thereof. 20 21 It would however be desirable to improve on the 22 solubility of known collagenase inhibitors and/or 23 stomelysin inhibitors (whether as the free base or the 24 salt) and, furthermore, increases in activity have also 25 It is not a simple matter, however, to been sought. 26 predict what variations in known compounds would be 27 desirable to increase or even retain activity; certain 28 modifications of known hydroxamic acid derivatives have 29 been found to lead to loss of activity. 30 31 According to a first aspect of the invention, there is 32 provided a compound of general formula I: 33

8

1
2
3
4
5 $R^{1}SO_{n}$ R^{3} R^{4} R^{5} R^{5} $R^{1}SO_{n}$ (I)

8 wherein:

9

10 R^1 represents a C_1 - C_6 alkyl, phenyl, thiophenyl, 11 substituted phenyl, phenyl(C_1 - C_6) alkyl, 12 heterocyclyl, (C_1 - C_6) alkylcarbonyl, phenacyl or 13 substituted phenacyl group; or, when n = 0, R^1 14 represents SR^X , wherein R^X represents a group:

15 16

17 18

19 20

21

22 R^2 represents a hydrogen atom or a C_1 - C_6 alkyl, C_1 - C_6 23 a l k e n y l , p h e n y l (C_1 - C_6) a l k y l , 24 cycloalkyl(C_1 - C_6) alkyl or cycloalkenyl(C_1 - C_6) alkyl 25 group;

26

27 R^3 represents an amino acid side chain or a C_1 - C_6 28 alkyl, benzyl, (C_1 - C_6 alkoxy)benzyl, 29 benzyloxy(C_1 - C_6 alkyl) or benzyloxybenzyl group;

3.0

R4 represents a hydrogen atom or a C₁-C₆ alkyl group;

32

33 R⁵ represents a hydrogen atom or a methyl group;

9 .

is an integer having the value 0, 1 or 2; and 1 n 2 represents a C1-C6 hydrocarbon chain, optionaly 3 A substituted with one or more C_1 - C_6 alkyl, phenyl 4 or substituted phenyl groups; 5 6 or a salt thereof. 7 8 Hereafter in this specification, the term "compound" 9 includes "salt" unless the context requires otherwise. 10 11 used herein the term "C1-C6 alkyl" refers to a 12 As straight or branched chain alkyl moiety having from 13 one to six carbon atoms, including for example, 14 methyl, ethyl, propyl, isopropyl, butyl, 15 pentyl and hexyl, and cognate terms (such as c^1-c^6 16 alkoxy") are to be construed accordingly. 17 18 The term "C1-C6 alkenyl" refers to a straight or 19 branched chain alkyl moiety having one to six carbons 20 and having in addition one double bond, of either E or 21 Z stereochemistry where applicable. This term would 22 include, for example, an alpha, beta-unsaturated 23 methylene group, vinyl, 1-propenyl, 1- and 2-butenyl 24 and 2-methyl-2-propenyl. 25 26 refers to a saturated "cycloalkyl" term The 27 alicyclic moiety having from 3 to 8 carbon atoms 28 and includes for example, cyclopropyl, cyclobutyl, 29 cyclopentyl and cyclohexyl. 30 31 32

WO 90/05719

1 The term "cycloalkenyl" refers to an unsaturated 2 alicycle having from 3 to 8 carbon atoms and includes cyclopropenyl, cyclobutenyl and cyclopentenyl, 3 4 cyclohexenyl. 5 The term "substituted", as applied to a phenyl or other 6 7 aromatic ring, means substituted with up to four substituents each of which independently may be C1-C6 8 alkyl, c_1-c_6 alkoxy, hydroxy, thiol, c_1-c_6 alkylthiol, amino, halo (including fluoro, chloro, bromo and iodo), 10 11 triflouromethyl or nitro. 12 13 The term "amino acid side chain" means a characteristic side chain attached to the -CH(NH₂)(COOH) moiety in the 14 15 following R or S amino acids: glycine, alanine, valine, 16 .. leucine, isoleucine, phenylalanine, tyrosine, 17 tryptophan, serine, threonine, cysteine, methionine, 18 asparagine, glutamine, lysine, histidine, arginine, 19 glutamic acid and aspartic acid. 20 21 The term "hydrocarbon chain" includes alkylene, 22 alkenylene and alkynylene chains of from 1 to 6 carbon 23 atoms. Preferably the carbon atom of the hydrocarbon 24 chain nearest to the hydroxamic acid group is a methylene carbon atom. 25 26 There are several chiral centres in the compounds 2.7 according to the invention because of the presence of 28 29 asymmetric carbon atoms. The presence of several 30 asymmetreic carbon atoms gives rise to a number of 31 diastereomers with the appropriate 32 stereochemistry at each chiral centre. General formula I and, where appropriate, all other formulae in this 33

PCT/GB89/01399

WO 90/05719

specification are to be understood to include all such 1 mixtures (for example racemic stereoisomers and 2 mixtures) thereof. Compounds in which the chiral centre adjacent the substituent R3 has S stereochemistry 4 and/or the chiral centre adjacent the substituent R^2 5 has R stereochemistry are preferred. 6 7 Further or other preferred compounds include those in 8 which, independently or in any combination: 9 10 R^1 represents a hydrogen atom or a C1-C4 alkyl, 11 phenyl, thiophenyl, benzyl, acetyl or benzoyl 12 13 group; 14 represents a C3-C6 alkyl (for example isobutyl) \mathbb{R}^2 15 group; 16 17 represents a benzyl or 4-(C1-C6)alkoxyphenylmethyl R^3 18 or benzyloxybenzyl group; 19 20 represents a C_1-C_4 alkyl (for example methyl) R^4 21 group; and 22 23 \mathbb{R}^5 represents a hydrogen atom. 24 25 Particularly preferred compounds include: 26 27 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenylthio-28 ı. methyl)-succinyl]-L-phenylalanine-N-methylamide, 29 30 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiophenyl-31 2. thio-methyl) succinyl]-L-phenylalanine-32 N-methylamide, 33

WO 90/05719

Τ.	3.	[4-(N-Hydroxyamino)-2R-Isobdcy1-35-(benzy1chio-
2		methyl) succinyl]-L-phenylalanine-N-methylamide,
3		
4	4.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(acetylthio-
5	·	methyl)succinyl]-L-phenylalanine-N-methylamide and
6		
7	5.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiolmethyl)
8	•	succinyl]-L-phenylalanine-N-methylamide
9		
10	6.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(benzoylthio-
11		methyl)succinyl]-L-phenylalanine-N-methylamide
12		
13	7.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(pivaloyl-
14		thiomethyl) succinyl]-L-phenylalanine-N-methyl-
15		amide
16		
17	8.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenyl-
18		thiomethyl) succinyl]-L-phenylalanine-N-methyl-
19	• .	amide sodium salt
20		
21	9.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-methoxy-
22		phenyl-thiomethyl) succinyl]-L-phenylalanine-N-
23		methylamide
24		
25	10.	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-hydroxy-
26		phenylthiomethyl) succinyl]-L-phenylalanine-N-
27		methylamide
28		
29	11	[4-(N-Hydroxyamino)-2R-isobutyl-3S-(2-thio-
30 "	•	phenethiomethyl)succinyl]-L-phenylalanine-N-
31		methylamide sodium salt
32	• •	
33	· .· -	
	•	

```
[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-methoxy-
 1
     12.
         phenylthiomethyl) succinyl]-L-phenylalanine-N-
 2
         methylamide sodium salt
 3
 4
         [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-tert-
 5
     13:
         butylphenylthiomethyl) succinyl]-L-phenylalanine-
 6
         N-methylamide
 7
 8
         [4-(N-Hydroxyamino)-2R-isobutyl-35-(2,4-di-
     14.
 9
         methylphenylthiomethyl)succinyl | -L-phenyl-
10 .
         alanine-N-methylamide
11
12
         bis-s, s'-{[4(N-Hydroxyamino-2R-isobutyl-
13
     15.
         3S-(thiomethyl) succinyl]-L-phenylalanine-N-methyl-
14
         amide } disulphide
15
16
         [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-bromo-
17
     16.
         phenylthio-methyl) succinyl]-L-phenylalanine-N-
18
         methylamide
19
20
         [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-chloro-
     17.
21
         phenylthiomethyl)succinyl]-L-phenylalanine-N-
22
         methylamide
23
24
          [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-methyl-
     18.
25
         phenylthiomethyl)succinyl]-L-phenylalanine-N-
26
27
         methylamide
28
          [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-acetyl)-
     19.
29
         aminophenylthiomethyl) succinyl]-L-phenylalanine-
30
         N-methylamide
31
32
33
```

14

20. [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenyl-1 sulphinylmethylsuccinyl]-L-phenylalanine-N-methyl-2 3 amide [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenyl-5 21. sulphonylmethylsuccinyl]-L-phenylalanine-N-methyl-6 7 amide 8 [4-(N-Hydroxyamino)-2R-isobutyl-3S-thiophenyl-9 . 22. sulphinylmethyl-succinyl]-L-phenylalanine-N-10 methylamide 11 12 23. [4-(N-Hydroxyamino)-2R-isobutyl-3S-thiophenyl-13 sulphonylmethyl-succinyl]-L-phenylalanine-N-14 methylamide 15 16 [4-(N-Hydroxyamino)-2R-isobuty1-3S-phenyl-17 24. sulphonylmethyl-succinyl]-L-phenylalanine-N-18 methylamide sodium salt 19 20 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(isobutyl-21 25. oxycarbonylamino) phenyl) thiomethyl-succinyl]-L-22 phenylalanine-N-methylamide 23 24 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-methyl-N-25 26. (tert-butoxycarbonyl)-glycylamino)phenyl)thio-26 methylsuccinyl]-L-phenylalanine-N-methylamide 27 28 and, where appropriate, their salts. Compounds 2 and 5 29 are especially preferred and compound 2 is the most 30 31 preferred, because of its good collagenase-inhibiting and protoglycanase-inhibiting activities. 32 33

PCT/GB89/01399 WO 90/05719

Compounds of general formula I may be prepared by any suitable method known in the art and/or by the following process, which itself forms part of the

invention.

According to a second aspect of the invention, there is provided a process for preparing a compound of general formula I as defined above, the process comprising:

(a) deprotecting a compound of general formula II

11
12
13
$$R^2$$
 N
 R^5
14
15
 $R^{1}SO$
 $R^{1}SO$
 R^{2}
 R^{3}
 R^{4}
 R^{5}
 R^{5}
(III)

wherein:

 R^{1} , R^{2} , R^{3} , R^{4} , R^{5} , A and n are as defined in general formula I and Z represents a protective group such as a benzyl group; or

(b) reacting a compound of general formula III

(III)

wherein:

(VIA)

 R^1 , R^2 , R^3 , R^4 , R^5 , A and n are as defined in general formula I,

with hydroxylamine or a salt thereof; or

6 (c) reacting a compound of general formula VIA

14 wherein

 \mathbb{R}^2 , \mathbb{R}^3 , \mathbb{R}^4 and \mathbb{R}^5 are as defined in general formula I,

either with a thiol of the general formula R¹S, wherein R¹ is as defined in general formula I to give a compound of general formula I in which A represents a methylene group and n is 0,

or with a cuprate of the general formula $(R^1S-A^1)_2CuLi$, wherein R^1 is as defined in general formula I and A^1 is such that $-A^1-CH_2$ — is identical to -A—, as defined in general formula I.

(d) optionally after step (a), step (b) or step (c) converting a compound of general formula I into another compound of general formula I.

Compounds of general formula I which are sulphoxides or sulphones can be derived from thiol compounds of general formula I by oxidation. Alternatively, thiols of general formula II or III may be oxidised. Compounds of general formula I which are disulphides (ie compounds wherein R1 represents SRX) may be derived from thiol esters of general formula I by milk oxidation, for example in air.

9 .

A compound of general formula II may be prepared from a compound of general formula III by reaction with an O-protected (such as benzyl) hydroxylamine. A compound of general formula III may be prepared by desterification (such as hydrolysis) of an ester of the general formula IV

wherein:

 R^1 , R^2 , R^3 , R^4 , R^5 , A and n are as defined in general formula I and R^6 represents C_1-C_6 alkyl, phenyl C_1-C_6 alkyl or substituted phenyl C_1-C_6 alkyl.

A compound of general formula IV can be prepared from an ester of general formula V or an acid of general formula VI

(VI)

formula I and R^6 represents c_1-c_6 alkyl, phenyl

 c_1-c_6 alkyl or substituted phenyl c_1-c_6 alkyl

by reaction with a thiol R^1SH , wherein R^1 is as defined

in general formula I, to give compounds wherein A

or by reaction with a cuprate of the general formula

(R¹S-A¹)₂CuLi, wherein R¹ is as defined in general

(V)

СООН

are as defined in general

1

3 4

5

6 7

8

9

wherein:

10

11

12

13

14

15

16

17

18

19

20

21

22

23 .

24 25

26

27 28

29 30

31 32 33

formula I and A^1 is such that $-A^1-CH_2-$ is identical to

-A-, as defined in general formula I.

represents a methylene group,

 R^2 , R^3 , R^4 and R^5

Esters of general formula V can be prepared by esterifying acids of general formula VI with an appropriate alcohol R⁶OH or other esterifying agent.

Compounds of general formula VIA can be prepared by reacting compounds of general formula VI with

hydroxylamine or a salt thereof.

An acid of general formula VI can be prepared by reacting a malonic acid derivative of general formula VII (VII) wherein: R^2 , R^3 , R^4 and R^5 are as defined in general formula I with formaldehyde in the presence of pyridine. An acid of general formula VII can in turn be prepared by desterifying (for example hydrolysing) a compound of general formula VIII (VIII) wherein: R^2 , R^3 , R^4 and R^5 are as defined in general formula I and R^6 represents C_1 - C_6 alkyl, phenyl C_1-C_6 alkyl or substituted phenyl C_1-C_6 alkyl.

A compound of general formula VIII can be prepared by reacting a compound of general formula IX with a compound of general formula X

R⁶O₃C COOH

:

(IX) (X)

CONR⁴R⁵

11 wherein:

 R^2 , R^3 , R^4 and R^5 are as defined in general formula I and R^6 represents C_1-C_6 alkyl, phenyl C_1-C_6 alkyl or substituted phenyl C_1-C_6 alkyl.

The starting materials and other reagents are either available commercially or can be synthesised by simple chemical procedures.

For example, a substituted acid of general formula IX may be prepared by reacting an ester of the general formula XI

26 R² CO₂1

. (XI)

wherein Y represents halo and R^5 is as defined above and R^2 and R^6 as defined above, with a malonate derivative of the general formula XII

$$R^6O_2C$$
 CO_2R^6 (XII)

21

wherein R^6 is as defined above with the proviso that 1 when R⁶ is aromatic in general formula XI it is 2 aliphatic in general formula XII or vice versa, and 3 selectively de-esterifying. 5 Compounds of general formula XI can simply be derived 6 from amino acids, which can be obtained 7 enantiomerically pure form, enabling a choice of 8 optically active compounds of general formula I to be 9 10 prepared. 11 Compounds of general formulae II and III are valuable 12 intermediates in the preparation of compounds of 13 general formula I. According to a third aspect of the 14 invention, there is therefore provided a compound of 15 general formula II. According to a fourth aspect of the 16 invention, there is provided a compound of general 17 formula III. 18 19 As mentioned above, compounds of general formula I are 20 useful in human or veterinary medicine as they are 21 active inhibitors, of metalloproteases involved in 22 tissue degradation. 23 24 According to a fifth aspect of the invention, there is 25 provided a compound of general formula I for use in 26 human or veterinary medicine, particularly in the 27 management (by which is meant treatment of prophylaxis) 28 of disease involving tissue degradation, in particular 29 rheumatoid arthritis, and/or in the promotion of wound 30 healing. 31

According to a sixth aspect of the invention, there is provided the use of a compound of general formula I in the preparation of an agent for the management of disease involving tissue degradation, particularly rheumatoid arthritis, and/or in the promotion of wound Compounds of general formula I can therefore be used in a method of treating disease involving tissue degradation, particularly rheumatoid arthritis, and/or in a method of promoting wound healing, the method in either case comprising administering to a human or animal patient an effective amount of a compound of general formula I.

16·

The potency of compounds of general formula I to act as inhibitors of collagenase (a metalloprotease involved in tissue degradation) was determined by the procedure of Cawston and Barrett, (Anal. Biochem., 99, 340-345, 1979) and their potency to act as inhibitors of stromelysin was determined using the procedure of Cawston et al (Biochem. J., 195, 159-165 1981), both of which techniques are to be described more fully in the examples and are incorporated by reference herein so far as the law allows.

According to a seventh aspect of the invention, there is provided a pharmaceutical or veterinary formulation comprising a compound of general formula I and a pharmaceutically and/or veterinarily acceptable carrier. One or more compounds of general formula I may be present in association with one or more non-toxic pharmaceutically and/or veterinarily acceptible carriers and/or diluents and/or adjuvents and if desired other active ingredients.

23

According to an eighth aspect of the invention, there is provided a process for the preparation of a pharmaceutical or veterinary formulation in accordance with the seventh aspect, the process comprising admixing a compound of general formula I and a pharmaceutically and/or veterinarily acceptable carrier.

8

10

11

12

13

14

Compounds of general formula I may be formulated for administration by any route and would depend on the disease being treated. The compositions may be in the form of tablets, capsules, powders, granules, lozenges, liquid or gel preparations, such as oral, topical, or sterile parental solutions or suspensions.

15 16

Tablets and capsules for oral administration may be in 17 unit dose presentation form, and may contain 18 conventional excipients such as binding agents, 19 example syrup, acacia, gelatin, sorbitol, tragacanth, 20 or polyvinyl-pyrollidone; fillers for example lactose, . 21 sugar, maize-starch, calcium phosphate, sorbitol or 22 glycine; tabletting lubricant, for example 23 magnesium sterate, talc, polyethylene glycol or 24 silica; disintegrants, for example potato starch, 25 agents such as sodium lauryl wetting acceptable 26 The tablets may be coated according to sulphate. 27 methods well known in normal pharmaceutical practice. 28 Oral liquid preparations may be in the form of, for 29 aqueous or oily suspensions, solutions, 30 emulsions, syrups or elixirs, or may be presented as a 31 for reconstitution with water or other dry product 32 suitable vehicle before use. Such liquid 33

24

preparations may contain coventional additives such as suspending agents, for example sorbitol, 2 cellulose, glucose syrup, gelatin, 3 hydrogenated edible fats; emulsifiying agents, for 4 sorbitan monooleate, or acacia; 5 example lecithin, non-aquieous vehicles (which may include edible oils). for example almond oil, fractionated coconut oil, oily esters such as glycerine, propylene glycol, 8 or ethyl alcohol; preservatives, for example methyl or 9 propyl p-hydroxybenzoate or sorbic acid, 10 desired conventional flavouring or colouring agents. 11

12 13

> 14 15

16

17

18

19

The dosage unit involved in oral administration may contain from about 1 to 250 mg, preferably from about 25 to 250 mg of a compound of general formula I. A suitable daily dose for a mammal may vary widely depending on the condition of the patient. However, a dose of a compound of general formula I of about 0.1 to 300mg/kg body weight, particularly from about 1 to 100 mg/kg body weight may be appropriate.

20 21 22

23

24

25

26

For topical application to the skin the drug may be made up into a cream, lotion or ointment. Cream or ointment formulations that may be used for the drug are conventional fomulations well known in the art, for example, as described in standard text books of pharmaceutics such as the British Pharmacopoeia.

27 28

For topical applications to the eye, the drug may be made up into a solution or suspension in a suitable sterile aqueous or non-aqueous vehicle. Additives, for instance buffers such as sodium metabisulphite or disodium edeate; preservatives including bactericidal

25

agents, such as phenyl mercuric and fungicidal 1 or nitrate, benzalkonium chloride 2 and thickening agents such as chlorohexidine, 3 hypromellose may also be included. 4 5 employed for the topical administration dosage 6 The depend on the size of the area being will, of course, 7 For the eyes each dose will be typically in 8 the range from 10 to 100 mg of the compound of general 9 formula I. 10 11 active ingredient may also be administered The 12 parenterally in a sterile medium. The drug 13 depending on the vehicle and concentration used, can 14 either be suspended or dissolved in the vehicle. 15 Advantageously, adjuvants such as a local anasthetic, 16 preservative and buffering agents can be dissolved in 17 the vehicle. 18 19 For use in the treatment of rheumatoid arthritis the 20 compounds of this invention can be administered by 21 the oral route or by injection intra-articularly into 22 The daily dosage the affected joint. for 23 mammal will be in the range of 10 mgs to 1 gram of a 24 compound of general formula I. 25 26 The following examples illustrate the invention, but 27 are not intended to limit the scope in any way. 28 following abbreviations have been used in 29 Examples:-30 31

20

21

22

· .23 . 24

- Dicyclohexylcarbodiimide 1 DCC DCM - Dichloromethane 2 DCU - Dicyclohexylurea 3 - Diisopropyl ether DIPE 4 - N, N-dimethylformamide 5 DMF - Hydroxybenztriazole HOBT 6 - N-Methylmorpholine 7 MMM - Trifluoroacetic acid 8 TFA - Tetrahydrofuran 9 THF WSCDI - N-(Dimethylaminoethyl)-N'-ethylcarbodiimide 10 11 12 Example 1 13 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenylthiomethyl)-14 succinyl]-L-phenylalanine-N-methylamide 15 16 17 18 19

a) 2R-Bromo-5-methylpentanoic acid.

PhS

(100g, 0.76 mol) and potassium bromide 25 D-Leucine (317.5g, 2.67 mol) were dissolved in aqueous acid 26 (150ml concentrated sulphuric acid in 500ml of water). 27 The solution was cooled to -20 and sodium nitrite 28 (69.6q, 0.95 mol in water) was added over 1h taking 29 care to maintain the temperature between -1 and -20. 30 After addition was complete the mixture was kept at 00 31 for a further hour, then DCM was added and the mixture 32 stirred for a few minutes. The layers were separated 33

CONHOH

```
and the ageous phase was washed with further portions
1
                             The combined organic layers
    of DCM (5 x 250ml).
2
    were dried over magnesium sulphate then the solvent
 3
    removed to give the acid as a pale yellow oil (123.1g,
    0.63 mol, 83%)
 5
 6
     [alpha]_D = +38.0^{\circ} (c = 2, methanol)
7
8 .
              (250 \text{ MHz}, \text{ CDCl}_3) 4.29 (1H, t, J= 6.5Hz,
9
    deltam
    BrCHCO_2H), 1.91 (2H, t, J= 7Hz, CHCH_2CH), 1.83 (1H, m,
10
    Me_2CH), and 0.94 (6H, 2xd, J= 7Hz, (CH_3)_2CH)
11
12
    b) tert-Butyl 2R-Bromo-5-methylpentanoate.
13
14
    2R-Bromo-5-methylpentanoic acid
                                        (123g,
                                                 0.63 mol)
15
    was dissolved in DCM (400ml) and the solution cooled
16
    to -40° while isobutene was condensed in to roughly
17
                          Maintaining the temperature at
    double the volume.
18
                                    acid (4ml) was added
    -40° concentrated sulphuric
19
                  When the addition was
                                             complete
20
    dropwise.
                was allowed to warm to room temperature
    reaction
21
                   The resultant solution was concentrated
    overnight.
22
    to half the volume by removing the solvent at reduced
23
    pressure, then the DCM was washed twice with an equal
24
    volume of 10% sodium bicarbonate solution. The organic
25
                         over magnesium sulphate and the
                 dried
26
     solvent removed under reduced pressure to leave the
27
    title compound as a yellow oil (148.0g, 0.59 mol, 94%).
28
29
     [alpha]_D = +23.0^{\circ} (c = 2, methanol)
30
31
32
33
```

WO 90/05719

33

```
(250 MHz, CDCl<sub>3</sub>) 4.18 (1H,
1
    deltan
                                             t, J = 6.5Hz,
    BrCHCO_2H), 1.89 (2H, m, CHCH_2CH), 1.78 (1H, m, Me_2CH),
2
3
    1.49 (9H, s, (CH_3)_3C) and 0.94 (6H, 2xd, J= 7Hz,
     (CH<sub>3</sub>)<sub>2</sub>CH)
4
5
    deltac
            (63.9 MHz, CDCl<sub>3</sub>) 167.0, 82.0, 46.3, 43.4,
6
7
    27.6, 26.3, 22.2, and 21.6.
8
9
     c) Benzyl (2-benzloxycarbonyl-3R-(tert-butoxycarbonyl)-
     5-methylhexanoate.
10
11
12
     Dibenzyl malonate (124.5g, 0.44 mol) was taken up in
13
    dry DMF and potassium tert-butoxide (49.2g, 0.44
    mol) was added portionwise with stirring and cooling.
14
    When a homogeneous solution had formed it was cooled to
15
     00 then tert-butyl-2R-bromo-5-methylpentanoate
16
    (110.0g, 0.44 mol) in DMF (200 ml) was added dropwise
17
               When addition was complete the reaction was
18
     transfered to a cold room at <50 and left for 4 days.
19
     The reaction mixture was partitioned between ethyl
20
21
                     saturated ammonium chloride then the
     aqueous layer extracted with further ethyl acetate
22
     (4x500ml), drying and solvent removal
23
                                                left an oil
     (228g) heavily contaminated with DMF.
24
                                               This oil was
     taken into ether (1 litre)
                                   and washed with brine
25
     (2x11) then the organic layer dried
26
                                                 (magnesium
     sulphate), solvent removed under reduced pressure to
27 .
28
     leave the desired material (179g) contaminated with a
     small amount of dibenzyl malonate.
29
30
     [alpha]_D = +22.5^O (c = 2, methanol)
31
32
```

PCT/GB89/01399 WO 90/05719

29

```
delta_{H} (250 MHz, CDCl_{3}) 7.40 - 7.25 (10H, m, Aromatic
1
    H), 5.14 (4H, 2xABq, CH_2Ph), 3.77 (1H, d, J= 10Hz,
2
    Bno<sub>2</sub>CCHCO<sub>2</sub>Bn), 3.09 (1H, dt,
                                             J=
3
    CH_2CH_{CO_2}tBu), 1.50 (3H, m, CH_2 + CH_{CO_2}tBu) 1.41 (9H, s,
4
    C(CH_3)_3) and 0.88 (6H, 2xd, J= 7Hz).
5
6
    d) [4-Benzyloxy-3-benzyloxycarbonyl-2R-isobutyl-
7
     succinyl]-L-phenylalanine-N-methylamide
8
9
     Benzyl(2-benzyloxycarbonyl-5-methyl-3R-tert-butoxycarb-
10
     onyl)-hexanoate (281.4g, 0.56 mol) was taken up in 5%
11
    water in TFA (410 ml) and allowed to stand at 5^{\circ}
12
                 After this time the TFA was evaporated
13
     under reduced pressure then the residue partitioned
14
     between DCM (11) and brine (200ml). Solvent removal
15
     left an oil which crystallised on standing (230g).
16
17
     The crude acid from this reaction was dissolved in DMF
18
     (11), then HOBT (95.3g, 0.64 mol), NMM (64g, 0.64 mol)
19
     and phenylalanine-N-methylamide (113.0g, 0.64 mol) were
20
                                    The mixture was cooled
     added at room temperature.
21
     to 0° before dropwise addition of DCC (131.0g, 0.64
22
     mol) in THF (11). This solution was stirred to room
23
     temperature over the weekend. The precipitated DCU was
24
     removed by filtration then the solvents were removed
25
     from the filtrate under reduced pressure to leave an
26
     oil. This oily residue was dissolved in ethyl acetate
27
                                               10% sodium
     then washed with 10% citric acid,
28
     bicarbonate and saturated brine. The organic layer was
29
     dried (magnesium sulphate), filtered then the solvent
30
     removed under reduced pressure to
                                            give the title
31
     compound as an oil (400g). This material was columned
```

on silica using gradient elution (0 -

50%

32

WO 90/05719

```
1
     acetate in hexane) to remove impurities
                                                and
                                                      separate
 2
        small amount of the minor diastereoisomer.
                                                           The
     material from the column (195g) was recrystallised
 3
            DIPE to give the title compound as a white
 4
 5
     crystalline solid (140.2g, 0.25 mol, 47%)
 6
     m.p. 98 -990
 7
     Analysis calculated for C33H38N2O6
 8
 9
     Requires C 70.95 H 6.86 N 5.01
10
     Found
              C 70.56 H 6.89 N 5.06
11
12
                       CDCl<sub>3</sub>) 7.42 - 7.13 (15H ,m, Aromatic
             (250MHz,
13
     H), 6.58 (1H,
                       d,
                             J=7.7Hz, CONH), 5.75 (1H,
     CONHMe), 5.20 - 5.05 (4H, m, OCH_2Ph), 4.50 (1H, dt, J=
14
15
     6.9,7.7Hz, CHCH<sub>2</sub>Ph), 3.79 (1H,
                                             d,
                                                   J=9.1Hz
     CH(CO_2Bn), 3.15 - 2.91 (2H, m, CH_2Ph), 2.65 (3H, d, J=
16
17
     4.8Hz, CONHC\underline{H}_3), 1.52 (1H, m, CHC\underline{H}_2CH), 1.32 (1H,
     C\underline{H}(CH_3)), 1.05 (1H, m, CHC\underline{H}_2CH), and 0.74 (6H, 2xd, J=
18
19
     6.5Hz, CH(CH_3)_2)
20
21
     e) [4-Hydroxy-2R-isobutyl-3-ethenylsuccinyl]-L-phenyl-
22
     alanine-N-methylamide.
23
     [4-Benzyloxy-3-benzyloxycarbonyl-2R-isobutylsuccinyl]-
24
25
     L-phenylalanine-N-methylamide (29.6g, 53mmol) was taken
     up in ethanol, ammonium formate (16.7g, 265mmol) added
26
27
     followed by 10%
                        palladium
                                     on
                                         charcoal (6g) as a
28
     slurry in isopropyl alcohol.
                                     After 30 minutes at room
29
     temperature the catalyst was removed by filtration,
30
    then washed with ethanol to give a solution
     crude diacid. To this was added piperidine (5.0g)
31
     the mixture stirred at room temperature for 15 minutes
32
33
     before
               addition
                           of
                                 aqueous formaldehyde (40%
```

```
After 18 hours at room temperature
                25ml).
     solution,
 1
                                               Solvents were
                    was refluxed for 1 h.
    the mixture
 2
                                pressure and the residue
     removed under reduced
 3
     partitioned between ethyl acetate
                                            and citric acid.
 4
    The acid layer was extracted with further portions
 5
    ethyl acetate (2x250ml), the combined organic layers
 6
           extracted with potassium carbonate (3x200ml).
 7
     These base extracts were acidified to pH 4 and
 8
     re-extracted with DCM then the organic layer dried
 9
                     magnesium sulphate. Solvent removal
10
     over
     under reduced pressure gave the desired product as a
11
     white solid (9.35g, 27.0mmol, 51%).
12
13
     m.p. 149-151°C
14
15
     delta_H (250MHz, CDCl_3) 8.37 (2H, d, J=9.0Hz, CONH),
16
                            7.27 - 7.06 (5H, m, Aromatic
     7.39 (1H, m, CONHMe),
17
     H), 6.40 (1H, s, C_{\underline{H}_2}CHCO_2H), 5.78 (1H, s, C_{\underline{H}_2}CHCO_2H),
18
     4.93 (1H, q, J= 7Hz, C\underline{H}CH_2Ph), 3.92 (1H, m, CH_2C\underline{H}CONH),
19
     2.95 (2H, m, C_{\underline{H}_2}Ph), 2.71 (3H, d, J=4.1Hz, NHC_{\underline{H}_3}),
20
     1.68 (1H, m), 1.45 (2H, m), and 0.86 (6H, 2xd, J=
21
     5.8Hz, CH(CH_3)_2).
22
23
     delta<sub>C</sub> (63.9Hz, CDCl<sub>3</sub>) 173.3, 172.8, 169.6, 139.1,
24
     136.3, 129.2, 128.3, 127.0, 126.6, 54.4, 43.5, 41.4,
25
     39.1, 26.2, 25.7, 22.5 and 22.4
26
27
     f) [4-Hydroxy-2R-isobutyl-3S-(phenylthiomethyl)-
28
     succinyl]-L-phenylalanine-N-methylamide
29
30
     [4-Hydroxy-2R-isobuty-3-ethenylsuccinyl]-L-phenyl-
31
     alanine-N-methylamide (15.0g, 44mmol) was dissolved in
32
     thiophenol
33
```

```
(150ml) and the mixture stirred in the dark under
     nitrogen at 600 for 2 days. Ether was added to the
2
     cooled reaction mixture and the precipitated product
3
     collected by filtration. The solid was washed with
     large volumes of ether and dried under vacuum to give
5
     the title compound (13.1g, 28.7mmol, 65%).
·6
7
     m.p. 199-201°C
 8
     Analysis calculated for C25H32N2O4S
 9
     Requires C 65.76 H 7.06 N 6.14 S 7.02
10
             C 65.69 H 7.06 N 6.07 S 7.05
11
12
     delta_{H} (250MHz, D_{6}-DMSO) 8.40 (1H, d, J= 9Hz, CON_{H}),
13
     7.82 (1H, m, CON_{HMe}), 7.35 - 7.10 (7H, m, Aromatic
14
     H), 7.04 (3H, m, Aromatic H), 4.62 (1H, m, CHCH<sub>2</sub>Ph),
15
16
     2.94 (1H, dd, J = 14,5Hz, CHCH_2Ph), 2.89
                                                   (1H, dd, J=
     14,9Hz, CHC\underline{H}_2Ph), 2.62 (3H, d, J= 4.5Hz, CONHC\underline{H}_3), 2.41
17
     (3H, m, 2xCH + CH<sub>2</sub>SPh), 2.23 (1H, d, J= 12Hz,
18
     1.43 (1H, m, CHC\underline{H}_2CH), 1.30 (1H, bm, C\underline{H}(CH<sub>3</sub>)<sub>2</sub>), 0.90
19
     (1H, m, CHC\underline{H}_2CH) and 0.78 (6H, 2xd, J= 6.5Hz, CH(C\underline{H}_3)<sub>2</sub>.
20
21
     g) [4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenylthio-
22.
     methyl) succinyl]-L-phenylalanine-N-methylamide
23
24
     [4-Hydroxy-2R-isobutyl-3S-(phenylthiomethyl)succinyl]-
     L-phenylalanine-N-methylamide (16.8g,
                                                  37 mmol) and
26
                        mmol) were dissolved in DCM / DMF
27
     HOBT (6.6g,
                   44
     (4:1) and the mixture cooled to 00 before adding WSCDI
28
     (8.5g, 44 mmol) and NMM (4.5g, 44 mmol).
                                                   The mixture
     was stirred at 00 for 1h to ensure complete formation
30
     of the activated ester. Hydroxylamine hydrochloride
31
     (3.8q, 55 mmol) and NMM (5.6g, 55 mmol) were dissolved
32
     in DMF then this mixture added dropwise to the cooled
3.3
```

1

```
solution of the activated ester. After 1h the reaction
     was poured into ether / water (1:1) whereupon the
     desired product precipitated as white crystals. These
 3
     were collected by filtration, further washed with ether
 4
     and water then dried under vacuum
                                                   at
                                                        50°.
 5
     material was recrystallised from methanol / water (1:1)
 6
     to remove a trace of the minor diastereomer (9.03g,
 7
     19.2 mmol, 52%).
 8
9
     m.p. 227-229°C
10
11
     [alpha]_D = -88^\circ (c = 10 , methanol)
12
13
     delta_{H} (250MHz, D_{6}-DMSO) 8.84 (1H, d, J= 1.5Hz, NHO\underline{H}),
14
     8.35 (1H, d, J= 8.7Hz, CONH), 7.87 (1H, m, CONHMe),
15
     7.29 - 6.92 (11H, m, Aromatic H + N\underline{H}OH), 4.60 (1H, m,
16
     C_{\underline{H}}CH_{2}Ph), 2.94 (1H, dd, J= 13.5,4.3, CHC_{\underline{H}_{2}}Ph), 2.77
17
     (1H, dd, J= 13.5,10, CHC\underline{H}_2Ph), 2.60 (3H, d,J= 4.6Hz),
18
     2.53 (1H, m), 2.41 (1H, m), 2.20 (1H, dd,
19
     13.4,2.2Hz, C_{\underline{H}_2}SPh), 2.09 (1H, dd, J=13.4,2.4Hz,
20
     C_{\underline{H}_2}SPh), 1.38 (2H, m, C_{\underline{H}}Me_2 + CHC_{\underline{H}_2}CH), 0.88 (1H,
21
     m, CHCH_2CH), 0.82 (3H, d, J=6.4Hz, CH(CH_3)_2), and 0.74
22
      (3H, d, J+ 6.4Hz, CH(CH<sub>3</sub>)<sub>2</sub>).
23
24
     delta<sub>C</sub> (63.9MHz, D<sub>6</sub>-DMSO) 172.9, 171.6, 166.3, 138.1,
25
     136.7, 129.1, 128.9, 128.0, 127.3, 126.4, 125.2, 54.2,
26
     46.4, 46.0, 37.7, 32.4, 25.6, 25.2, 24.2, and 21.7.
27
28
29
30
31
32
33
```

Example 2

1. 2 3

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiophenylthiometh-yl) succinyl]-L-phenylalanine-N-methylamide

4 5 6

7 8 9

12 13 14

10 11

a) [4-N-Hydroxy-2R-isobutyl-3S-(thiophenylthiomethyl) succinyl]-L-phenylalanine-N-methylamide

15 16

title compound prepared 17 The was [4-Hydroxy-2R-isobutyl-3-ethenylsuccinyl]-L-phenyl-18 (400mg, 1.16mmol) by the method alanine-N-methylamide 19 described in example 1f, substituting thiophenethiol in 20 the place of thiophenol to give a material (320mg, 21 22 0.73mmol, 63%) with the following characteristics.

23

24 m.p. 184-186°C

25

delta_H (250MHz, D_6 -DMSO) 8.29 (1H, d, J= 8.1Hz, CONH), 26 CONHMe), 7.57 27 7.84 (1H, m, (1H, d, J= 5.1Hz,Thiophene H), 5H, m, Aromatic H), 7.00 28 Thiophene H), 4.50 (1H, m, CHCH2Ph), 2.91 (1H, 29 $CHCH_2Ph$), 2.75 (1H, m, $CHCH_2Ph$), 2.56 (3H, 30 4.0Hz, CONHCH₃), 2.34 (3H, m), 1.99 (1H, d, J= 9.3Hz, 31

32

WO 90/05719 PCT/GB89/01399

```
CH_2SHet), 1.42 (1H, m, CHCH_2CH), 1.29 (1H, bm,
1
     C_{H}(CH_3)_2, 0.87 (1H, m, CHC_{H_2}CH), 0.79 (3H, d, J=
2
     6.4Hz, CH(C\underline{H}_3)_2), and 0.72 (3H, d, J= 6.4Hz, CH(C\underline{H}_3)_2).
3
 4
     b) [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiophenylthio-
5
     methyl) succinyl]-L-phenylalanine-N-methylamide
 6
 7
     Prepared by the method described in example 1g to
 8
     give material with the following characteristics
9
10
     m.p. 236-238°C
11
12
     Analysis calculated for C23H30N2O4S2
13
     Requires C 57.84 H 6.54 N 8.80
14
               C 57.64 H 6.48 N 8.85
15
     Found
16
     delta<sub>H</sub> (250MHz, D<sub>6</sub>-DMSO) 8.80 (1H, s, CONHO<u>H</u>), 8.08
17
     (1H, d, J=8Hz, CONH), 7.52 (1H, m, CONHMe), 7.32 (1H,
18
     dd, J = 4.6, 2.9 Hz, Thiophene H), 7.17 - 6.95 (5H, m,
19
     Aromatic H), 6.89 (2H, m, Thiophene H), 4.46
20
     m, CHCH_2Ph), 2.89 (1H, dd, J=13.6,4.4Hz, CHCH_2Ph), 2.72
21
     (1H, dd, J= 13.6, 10.5Hz, CHCH<sub>2</sub>Ph), 2.54
22
     4.3Hz, CONHC\underline{H}_3), 2.46 (1H, d, J= 12.1Hz, C\underline{H}_2S),
23
     (1H, bt, J= 10.2Hz), 2.14 (1H, bt, J= 10.2Hz), 1.98
24
     (1H, dd, J=12.7,2.5Hz, CHC\underline{\text{H}}_2Ph), 1.35 (1H, bt, J=
25
     11.4Hz, CHC\underline{H}_2CH), 1.22 (1H, bm, CH(C\underline{H}_3)_2), 0.86 (1H,
26
     bt, J=12.6Hz, CHCH_2CH), 0.74 (3H, d, J=6.3Hz,
27
     CH(CH_3)_2, and 0.68 (3H, d, J= 6.4Hz, CH(CH_3)_2).
28
29
     delta<sub>C</sub> (63.9MHz, D<sub>6</sub>-DMSO) 172.5, 171.6, 166.1, 138.0,
30
     133.8, 132.7, 129.4, 129.2, 128.1, 127.8, 126.5, 54.2,
31
     46.2, 46.0, 38.5, 37.6, 25.8, 25.2, 24.2, and 21.7.
33
```

Example 3 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(benzylthiomethyl) succinyl]-L-phenylalanine-N-methylamide соинон PhCH₂S Prepared by the method described in example 1g to give material with the following characteristics m.p. Analysis calculated for C27H37N3O5S.0.5H2O Requires C 61.81 H 7.30 N 8.00 19 . Found C 61.85 H 7.15 N 7.45 delta_H (250MHz, D_6 -DMSO) 8.40 (1H, s, CONHO<u>H</u>), 8.22 (1H, m, NHMe), 7.20 (5H, m, Aromatic H), 6.58 (4H, m), 4.10 (1H, m, CHC \underline{H}_2 Ph), 3.22 (3H, s, OC \underline{H}_3), 3.04 - 2.45 (4H, m, $2xCH_2Ar$), 2.42 (3H, d, J= 6Hz, $NHCH_3$), 2.32 -2.08 (4H, m), 0.78 (2H, m, $CHC_{\frac{H}{2}}CH$), and 0.40 - 0.18 $(7H, m, (CH_3)_2CH)$

```
1
     Example 4
 2
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(acetylthiomethyl)
 3
     succinyl]-L-phenylalanine-N-methylamide
 4
 5
 б
 7
 8
 9
10
11
12
     Prepared by the method described in example 1g to
13
     give material with the following characteristics
14
15
     m.p. 226-227°C
16
17
     Analysis calculated for C21H31N3O5S.H2O
18
     Requires C 55.37 H 7.30 N 9.22
19
     Found
                C 55.57 H 6.99 N 9.53
20
21
     delta_{H} (250MHz, D_{6}-DMSO) 8.84 (1H, s, NHO\underline{H}), 8.36 (1H,
22
     d, J = 8Hz, CONH), 7.80 (1H, d, J = 6Hz, NHMe), 7.20 (%h,
23
     m, Aromatic H), 4.58 (1H, m, CHCH2Ph), 3.16 - 2.62
24
      (2H, m, CHC_{H_2}Ph), 2.54 (3H, d, J= 4Hz, NHC_{H_3}), 2.22
25
      (3H, s, CH_3COS), 2.36 - 2.10 (4H, m, CHCHCH_2S), 1.36
26
      (2H, m, CHC\underline{\text{H}}_2CH), and 0.98 - 0.66 (7H, m, C\underline{\text{H}}(C\underline{\text{H}}_3)<sub>2</sub>).
27
28
29
30
31
32
33
```

```
1
    Example 5
2
3
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiolmethyl)
    succinyl]-L-phenylalanine-N-methylamide
4
5
6
7
8
                              CONHOH
10
11
12
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(acetylthiomethyl)
13
    succinyl]-L-phenylalanine-N-methylamide (30mg,
     0.06mmol) was stirred
                                in methanol (3ml) with
14
    methylamine (1ml methanolic solution)
15
                                                 at
                    After 30 minutes the crystalline
     temperature.
16
17
     product (20mg, 0.05mmol, 74%) was filtered off and
    dried.
18
19
    m.p. 234°C
20
    Analysis calculated for C19H39N3O4S.1.5H2O
21
22
     Requires C 54.10 H 7.63 N 9.94 S 7.60
            C 54.28 H 7.16 N 10.43 S 7.80
23
     Found
24
     delta_{H} (250MHz, D_{6}-DMSO) 8.28 (1H, d, J= 9Hz, NHOH),
25
     7.80 (1H, m, NHMe), 7.22 (5H, m, Aromatic H), 4.60 (1H,
26
27
     m, CHCH_2Ph), 3.08 - 2.56 (2H, m, CHCH_2Ph), 2.50 (3H, d,
     J=4Hz, NHCH_3), 2.40 - 2.02 (4H, m, CHCHCH_2SH), 1.44
28
     - 1.22 (2H, m, CHC\underline{H}_2CH) and 0.98 - 0.72 (7H, m,
29
30
     CH(CH_3)_2.
31
32
33
```

PCT/GB89/01399

31

32 33

Example 6 1 2 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(benzoylthiomethyl)-3 succinyl]-L-phenylalanine-N-methylamide 4 5 6 7 8 9 10 11 12 The title compound was prepared by the method described 13 in Example 1g to give material with the following 14 characteristics 15 16 m.p. 227 - 228⁰ 17 Analysis calculated for C21H31N3O5S 18 Requires C 62.50 H 6.66 N 8.41 19 C 62.32 H 6.67 N 8.40 20 Found 21 delta_H (250 MHz, CDCl₃:D₆DMSO (1:1)) 8.82 (1H, 22 NHOH), 8.25 (1H, d, J=8.4Hz, NHOH), 7.87 (2H, dd, 23 J=8.5, 1.1Hz), 7.60 (2H, m, Ar-H and CONH), 7.50 (2H, 24 t, J=8.2Hz), 7.28 (2H, d, J=8.4Hz), 7.16 (2H, t, 25 J=7.2Hz), 7.04 (1H, t, J=8.5Hz), 4.65 (1H, m, $C\underline{H}CH_2Ph$), 26 3.06 (1H, dd, J=14.1, 5.0Hz, $CHCH_2Ph$), 2.90 (1H, dd, 27 J=13.9, 10Hz, $CHC\underline{H}_2Ph$), 2.73 (2H, m $SC\underline{H}_2Ph$), 2.65 (3H, 28 d, J=4.7Hz, NHMe), 2.33 (1H, dt, J=11.0, 4.7Hz), 1.51 29 (1H, t, J=7Hz, CH_2CHMe_2), 1.24 (1H, m, $CHMe_2$), 0.97 30

(1H, t, J=7Hz, $C_{\underline{H}_2}$ CHMe₂), 0.84 (3H, d, J=6.5Hz, $C_{\underline{H}_2}$)

and 0.79 (3H, d; J=6.5Hz, $CHMe_2$).

Example 7

1 2 3

4.

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(pivaloylthiomethyl) succinyl]-L-phenylalanine-N-methylamide

5 6 7

8

10 11 12 H CONHOH NHMe

13 14

15 [4-Hydroxy-2R-isobutyl-3S-(pivaloylthiomethyl) succinyl]-L-phenylalanine-N-methylamide (0,8g, 1.7 16 mmol) and HOBT (0.31g, 2.1 mmol) were dissolved in 1:1 17 DCM/DMF and the mixture cooled to 0°C before adding 18 WSDCI (0.4g, 2.1mmol) and NMM (0.21g, 2.1mmol). The 19. mixture was stirred at 0°C for 1h to ensure complete 20 21 formation of the activated ester. Hydroxylamine hydrochloride (0.18g, 2.6mmol) and NMM (0.26g, 2.6mmol) 22 were dissolved in DMF then this mixture was added 23 dropwise to the cooled solution of the activated ester. 24 After 1h the reaction was poured into ether/water (1:1) 25 whereupon the desired product precipitated as white 26 crystals. These were collected by filtration, further 27 28 washed with ether and water, then dried under vacuum at 50°C . This material was recrystallised from 29 methanol/water (1:1) to remove a trace of the minor 30 diastereomer (0.38g, 0.7mmol, 45%). 31

32

33 m.p. 225°C

PCT/GB89/01399

```
[alpha]_D = -3.5^{\circ} (c=2, methanol)
 2
 3
     Analysis calculated for C24H39N3O5S.0.5 H2O
     Requires: C58.99 H7.84 N8.60
     Found:
               C58.96 H7.63 N9.55
 5
 6
 7
    delta_{H} (250MHz, D_{6}-DMSO) 8.81 (1H, s, J = 1.5Hz, NHOH),
    8.30 (1H, d, J=8Hz, CONH), 7.78 (1H, d, J=6Hz, CONHMe),
. 8
    7.27-7.03 (5H, m, aromatic H), 4.54 (1H, m, CHCH<sub>2</sub>Ph),
    2.94 (1H, dd, J = 12,5Hz, CHCH_2Ph), 2.79 (1H, dd, J =
10
    13,10Hz, CHCH_2Ph) 2.56 (3H, d, J = 4.5Hz, NHCH_3), 2.44
11
    (2H, m), 2.20 (1H, dd, J = 13,3Hz, CH<sub>2</sub>S), 2.07 (1H, dd)
12
    dt), 1.36 (2H, m), 1.13 (9H, s, C(CH_3)_3), 0.87 (1H, m,
13
    CH_2CH(CH_3)_2, 0.79 (3H, d, J = 6Hz, CH(CH_3)_2), and 0.74
14
     (3H, d, J = 6Hz, CH(CH<sub>3</sub>)<sub>2</sub>).
15
16
             (63.9MHz, D<sub>6</sub>-DMSO) 172.55, 171.59, 168.24,
17
    138.03, 129.18, 128.00, 126.24, 54.21, 46.48, 45.84,
18
    45.55, 37.61, 28.30, 27.13, 25.64, 25.25, 24.24, and
19
    21.63.
20
21
    Example 8
22
23
    [4-(N-Hydroxyamino)-2R-isobuty1-3S-(phenylthiomethy1)
24
    succinyl]-L-phenylalanine-N-methylamide sodium salt
25
26
27
28
29
30
31
32
33
```

```
[4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenylthiomethyl)
2 succinyl]-L-phenylalanine-N-methylamide (0,2g, 0.4
3 mmol) was dissolved in 20ml of methanol and 1eq of 0.1N
4 NaOH(aq) added. The solvent was removed in vacuo and
5 the residue dissolved in water and freeze-dried
6 (0.21g,0.4 mmol,100%).
```

m.p. 184°C

10 [alpha]_D = -7.7° (c=2, methanol)

11
12 delta_H (250MHz, D₆-DMSO) 8.62 (1H, s, J = 1.5Hz, NHO<u>H</u>),
13 8.28 (1H, d, J = 8Hz, CON<u>H</u>), 7.26 - 7.04 (10H, m,
14 aromatic H), 4.43 (1H, m, CHCH₂Ph), 3.00 (1H, dd, J =
15 14,4Hz, CHCH₂Ph), 2.84 (1H, dd, J = 14,10Hz, CHCH₂Ph),
16 2.55 (3H, d, J = 4.5Hz, NHCH₃), 2.46 (3H, m), 2.21 (1H,
17 m), 1.39 (1H, m), 1.14 (1H, m), 1.00 (1H,m), and 0.70
18

Example 9

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-methoxyphenyl-thiomethyl)

```
succinyl]-L-phenylalanine-N-methylamide[4-Hydroxy-2R-
    isobuty1-3S-(4-methoxyphenylthiomethyl)succinyl]-L-
    phenylalanine-N-methylamide (0,5g, 1 mmol) and HOBT
    (0.18g, 1.2 mmol) were dissolved in 1:1 DCM/DMF and the
 4
    mixture cooled to 0°C before adding WSDCI (0.23g,
 5
    1.2mmol) and NMM (0.12g, 1.2mmol). The mixture was
 6
    stirred at 0°C for 1h to ensure complete formation of
 7
    the activated ester. Hydroxylamine hydrochloride (0.1g,
 8
    1.5mmol) and NMM (0.15g, 1.5mmol) were dissolved in DMF
 9
    then this mixture was added dropwise to the cooled
10
    solution of the activated ester. After 1h the reaction
11
    was poured into ether/water (1:1) whereupon the desired
12
    product precipitated as white crystals. These were
13
    collected by filtration, further washed with ether and
14
    water, then dried under vacuum at 50°C. This material
15
    was recrystallised from methanol/water (1:1) to remove
16
    a trace of the minor diastereomer (0.36g, 0.7mmol,
17
    72%).
18
19
    m.p. 225°C
20
21
    [alpha]_D = +8^O (c=0.5, methanol)
22
23
    Analysis calculated for C26H35N3O5S
24
    Requires: C62.25 H7.04 N8.38
25
              C62.43 H7.09 N8.37
    Found:
26
27
    delta_{H} (250MHz, D_{6}-DMSO) 8.83 (1H, s, J = 1.5Hz, NHO_{H}),
28
    8.28 (1H, d, J = 8Hz, CONH), 7.83 (1H, d, J = 6Hz,
29
    CONHMe), 7.28 - 6.86 (9H, m, aromatic H), 4.52 (1H, m,
30
    CHCH_2Ph), 3.73 (3H, s, OCH3), 2.91 (1H, dd, J = 14,4Hz,
31
    CHCH_2Ph), 2.75 (1H, dd, J = 14,10Hz, CHC\underline{H}_2Ph), 2.57
32
    (3H, d, J = 4.5Hz, NHCH<sub>3</sub>), 2.50 - 2.34 (2H,m), 2.16 -
```

1 1.99 (2H, m, $CH_2CH(CH3)_2$) 1.36 (2H, m), 0.88 (1H, m, $CH_2CH(CH_3)_2$), 0.80 (3H, d, J = 6Hz, $CH(CH_3)_2$), and 0.73 (3H, d, J = 6Hz, $CH(CH_3)_2$).

5 delta_C (63.9MHz, D₆-DMSO) 172.79, 171.62, 168.39,
6 138.14, 131.34, 129.19, 128.00, 126.44, 114.59, 55.32,
7 54.20, 38.68, 25.63, 25.17, 24.26, and 21.70.

Example 10

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-hydroxyphenyl-thiomethyl) succinyl]-L-phenylalanine-N-methylamide

31 ·

[4-Hydroxy-2R-isobutyl-3S-(4-hydroxyphenylthiomethyl) succinyl]-L-phenylalanine-N-methylamide (0,4g, 0.8 mmol) and HOBT (0.15g, 1.0 mmol) were dissolved in 1:1 DCM/DMF and the mixture cooled to 0°C before adding WSDCI (0.20g, 1.0mmol) and NMM (0.1g, 1.0mmol). The mixture was stirred at 0°C for 1h to ensure complete formation of the activated ester. Hydroxylamine hydrochloride (0.09g, 1.3mmol) and NMM (0.13g,1.3mmol) were dissolved in DMF then this mixture was added dropwise to the cooled solution of the activated ester. After 1h the reaction was poured into ether/water (1:1)

WO 90/05719 PCT/GB89/01399

45

whereupon the desired product precipitated as white

```
2 crystals. These were collected by filtration, further
    washed with ether and water, then dried under vacuum at
 4 50°C. This material was recrystallised from
    methanol/water (1:1) to remove a trace of the minor
    diastereomer (0.13g, 0.2mmol, 31%).
 7
    m.p. 216°C
 8
 9
    [alpha]_D = -65^{\circ} (c=0.5, methanol)
10
11
12
    Analysis calculated for C25H33N3O5S
    Requires: C61.58 H6.82 N8.62
13
    Found:
               C61.43 H6.81 N8.08
14
15
    delta_{H} (250MHz, D_{6}-DMSO) 8.82 (1H, s, J = 1.5Hz, NHOH),
16
17
    8.26 (1H, d, J = 8Hz, CONH), 7.81 (1H, d, J = 6Hz,
    CONHMe), 7.27 - 6.64 (9H, m, aromatic H), 4.49 (1H, m,
18
    C\underline{H}CH_2Ph), 2.90 (1H, dd, J=14,4Hz, CHC\underline{H}_2Ph), 2.74 (1H,
    dd, J=14,10Hz, CHCH_2Ph), 2.57 (3H, d, J=4.5Hz,
20
    NHCH_3), 2.54 - 2.29 (2H, m), 2.14 - 1.98 (2H, m,
21
22
    CH_2CH(CH3)_2), 1.35 (2H, m), 0.88 (1H, m, CH_2CH(CH_3)_2),
    0.80 (3H, d, J = 6Hz, CH(CH<sub>3</sub>)<sub>2</sub>), and 0.73 (3H, d, J =
23
    6Hz, CH(CH<sub>3</sub>)<sub>2</sub>).
24
25
            (63.9MHz, D<sub>6</sub>-DMSO) 172.81, 171.66, 168.46,
26
    deltac
    156.50, 133.02, 132.17, 129.17, 128.02, 126.44, 124.17,
27
    116.00, 54.20, 46.35, 46.13, 37.59, 35.40, 25.62,
28
    25.16, 24.27, and 21.69.
29
30
31
32
33
```

```
Example 11
 2
    [4-(N-Hydroxyamino)-2R-isobutyl-3S-(2-thiophenethio-
 3
    methyl)succinyl]-L-phenylalanine-N-methylamide sodium
 4
    salt
 5
 6
 7
 8
 9
10
11
                            CONHONa
12
13
14
15
    [4-Hydroxyamino)-2R-isobutyl-3S-(2-thiophenethiomethyl)
16
    succinyl]-L-phenylalanine-N-methylamide (0,2g,
17
    mmol) was dissolved in 20ml of methanol and 1eq of 0.1N
18
    NaOH(aq) added. The solvent was removed in vacuo and
19
    the residue dissolved in water and freeze-dried
20
    (0.21g, 0.4 mmol, 100%).
21
22
    m.p. 170°C
23
24
    [alpha]_D = -67^O (c=1, methanol)
25
2.6
    delta_{H} (250MHz, d_{6}-DMSO), 7.51 (1H, d), 7.19 - 6.97
27
    (8H, m, aromatic H), 4.32 (1H, m, CHCH<sub>2</sub>Ph), 3.00 (1H,
28
    dd, J = 14,4Hz, CHCH_2Ph), 2.84 (1H, dd, J = 14,10Hz,
29
    CHCH_2Ph) 2.53 (3H, d, J = 4.5Hz, NHCH_3), 2.46 2.19 (3H,
30
    m), 1.37 (1H, m), 1.09 (1H, m), 0.93 (1H, m), and 0.67
31
    (6H, m)
32
```

PCT/GB89/01399

```
Example 12
```

[4-(N-Hydroxyamino)-2R-isobuty1-3S-(4-methoxyphenylthiomethyl)succinyl]-L-phenylalanine-N-methylamide sodium salt

WO 90/05719

[4-Hydroxyamino)-2R-isobutyl-3S-(4-methoxyphenylthio-methyl)succinyl]-L-phenylalanine-N-methylamide (0,1g, 0.2 mmol) was dissolved in 20ml of methanol and 1eg of 0.1N NaOH(aq) added. The solvent was removed in vacuo and the residue dissolved in water and freeze-dried (0.1g,0.2 mmol,100%).

m.p. 174°C

 $[alpha]_D = -58^{\circ}$ (c=1, methanol)

 $delta_H$ (250MHz, D_6 -DMSO _7.26 - 7.04 (10H, m, aromatic H), 4.31 (1H, m, $CHCH_2Ph$), 3.73 (3H, s, OCH_3), 3.25 -2.72 (2H, m, CHCH₂Ph), 2.50 (3H, s, NHCH₃), 2.36 (1H, m), 2.15 (1H, m), 1.37 (1H, m), 0.95 (1H, m), and 0.69 (6H, d, $CHCH_2(CH_3)_2$).

Example 13

2

4

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-tertbutylphenyl-thiomethyl) succinyl]-L-phenylalanine-N-methylamide

5

7 8 9 H CONHOH

13 14

10 11 12

15 16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32 33

[4-Hydroxy-2R-isobutyl-3S-(4-tertbutylphenylthiomethyl) succinyl]-L-phenylalanine-N-methylamide (5.0g, 10 mmol) and HOBT (1.76g, 12 mmol) were dissolved in 1:1 DCM/DMF and the mixture cooled to 0°C before adding WSDCI (2.3g, 12mmol) and NMM (1.2g, 12mmol). The mixture was stirred at 0°C for 1h to ensure complete formation of the activated ester. Hydroxylamine hydrochloride (1.0g, 15mmol) and NMM (1.2g, 15mmol) were dissolved in DMF then this mixture was added dropwise to the cooled solution of the activated ester. After 1h the reaction was poured into ether/water (1:1) whereupon the desired product precipitated as white crystals. These were collected by filtration, further washed with ether and water, then dried under vacuum at 50°C. This material was repeatedly recrystallised from methanol/water (1:1) to remove a trace of the minor diastereomer (0.7g, 1.3mmol, 14%).

```
M.p. 188.5 - 190^{\circ}C
 1
 2
    Analysis calculated for C29H41N3O4S
 3
    Requires: C66.00 H7.83 N7.96
 4
    Found:
                C65.80 H7.81 N7.76
 5
 6
    delta<sub>H</sub> (250MHz, D_6-DMSO) 8.83 (1H, s, NHO<u>H</u>), 8.33 (1H,
 7
    d, J = 8Hz, CONH), 7.86 (1H, d, J = 6Hz, CONHMe), 7.28
    - 6.90 (9H, m, aromatic H), 4.60 (1H, m, CHCH<sub>2</sub>Ph), 2.94
 9
    (1H, dd, J = 14,4Hz, CHCH<sub>2</sub>Ph), 2.77 (1H, dd, J =
10
    14,10Hz, CHC\underline{H}_2Ph), 2.58 (3H, d, J = 4.5Hz, NHC\underline{H}_3), 2.55
11
    -2.37 (2H, m), 2.22 - 2.08 (2H, m, CH_2CH(CH3)_2), 1.37
12
           m), 1.26 (9H, s, C(CH_3)_3), 0.88 (1H,
13
    C_{H_2}CH(C_{H_3})_2, 0.81 (3H, d, J = 6Hz, CH(C_{H_3})_2), and 0.74
14
     (3H, d, J = 6Hz, CH(CH<sub>3</sub>)<sub>2</sub>).
15
16
              (63.9MHz, D<sub>6</sub>-DMSO) 172.88, 171.59, 168.34,
    deltac
17
    147.87, 138.10, 133.09, 129.13, 127.95, 127.45, 126.36,
18
    125.70, 54.19, 54.20, 46.38, 46.06, 37.70, 34.20, 32.79
19
    31.24, 25.64, 25.19, 24.25, and 21.72.
20
21
    Example 14
22
23
    [4-(N-Hydroxyamino)-2R-isobuty1-3S-(2,4-
24
    dimethylphenylthiomethyl) succinyl]-L-phenylalanine-N-
25
    methylamide
26
27
28
29
30
31
32
33
```

[4-Hydroxy-2R-isobuty1-3S-(2,4-dimethylphenylthio-2 methyl)succinyl]-L-phenylalanine-N-methylamide (1.8g, 3.7 mmol) and HOBT (0.67g, 12 mmol) were dissolved in 4 1:1 DCM/DMF and the mixture cooled to 0°C before adding 5 WSDCI (0.86g, 4.5mmol) and NMM (0.45g, 4.5mmol). The 6 mixture was stirred at 0°C for 1h to ensure complete 7 formation of the activated ester. Hydroxylamine 8. hydrochloride (0.39g, 5.6mmol) and NMM (0.56g, 5.6mmol) 9 were dissolved in DMF then this mixture was added 10 dropwise to the cooled solution of the activated ester. 11 After 1h the reaction was poured into ether/water (1:1) 12 whereupon the desired product precipitated as white 13 crystals. These were collected by filtration, further 14 washed with ether and water, then dried under vacuum at 15 50°C. This material was repeatedly recrystallised from 16 methanol/water (1:1) to remove a trace of the minor 17 diastereomer (1.08g, 2.2mmol, 58%). 18

19

m.p. 226°C (dec.)

20

Analysis calculated for C₂₇H₃7N₃O₄S
Requires: C64.90 H7.46 N8.41

23 Found: C65.15 H7.48 N8.40

24

25 delta_H (250MHz, D₆-DMSO) 8.83 (1H, s, NHOH), 8.32 (1H, 26 d, J = 8Hz, CONH), 7.85 (1H, d, J = 6Hz, CONHMe), 7.30 27 - 6.71 (9H, m, aromatic H), 4.56 (1H, m, CHCH2Ph), 2.91 28 (1H, dd, J = 14,4Hz, CHCH₂Ph), 2.76 (1H, dd, J =29 14,10Hz, CHC \underline{H}_2 Ph), 2.57 (3H, d, J = 4.5Hz, NHC \underline{H}_3), 2.53 30 - 2.38 (2H, m), 2.23 (3H, s, $C_6H_5(CH_3)$ 2), 2.13 (3H, s, 31 $C_6H_5(CH_3)$, 1.30 (2H, m), 0.89 (1H, m, $CH_2CH(CH_3)_2$), 32 0.81 (3H, d, J = 6Hz, CH(CH₃)₂), and 0.74 (3H, d, J =33 6Hz, $CH(C\underline{H}_3)_2$).

2

Example 15

8

9 10

NHMe H// CONHOH CONHOH Н NHMe

11 12

13

[4(N-Hydroxyamino-2R-isobutyl-3S-(acetylthiomethyl) 14 succinyl]-L-phenylalanine-N-methylamide (1.0g, 2.4 15 mmol) was dissolved in 750ml methanol and 350ml pH 7 16 Left to stand overnight and solvent buffer added.

17 removed in vacuo to 2/3 volume, left to crystallise for 18 a further two hours. Filtered and dried to give 0.87g

19 off-white crystals

20 21

Analysis calculated for C38H56N6O8S2.1.9H2O

22 Requires: C55.34 H6.93 N9.88

23 C55.44 H7.32 N10.21 Found:

24 25

Example 16

26 27

28

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-bromophenylthiomethyl) succinyl]-L-phenylalanine-N-methylamide

29 30 31

32

```
1
      Prepared by the method described in example 1g to give
  2
      material with the following characteristics.
      m.p. 225 -229°C
  5
  6
      [alpha]_D = -164.8^{\circ}
  7
  8
      Analysis calculated for c_2^5 H_{32} Br N_3 O_4 S
      Requires: C54.40 H5.89 N7.40
 10
      Found:
                  C54.54 H5.86 N7.63
 11
 12
      delta<sub>H</sub> (250MHz, D_6-DMSO) 8.83 (1H, s, NHO<u>H</u>), 8.35 (1H,
 13
      d, J = 8Hz, CONH, 7.90 (1H, q, J = 6Hz, CONHMe), 7.35
 14
      - 6.87 (9H, m, aromatic H), 4.64 (1H, m, CHCH<sub>2</sub>Ph), 2.94
 15
      (1H, dd, J = 14,4Hz, CHCH<sub>2</sub>Ph), 2.76 (1H, t, J = 13Hz)
 16
      CHC\underline{H}_2Ph) 2.60 (3H, d, J = 5Hz, NHC\underline{H}_3), 2.55 - 2.35 (2H,
 17
      m, CH_2S), 2.15 (1H, t, J = 10Hz, CHCO), 2.01 (1H, d, J
 18
      = 11.5 Hz, CHCO),
                               1.37 (2H, m),
                                                     0.88 (1H,
 19
      C_{\underline{H}_2}^2CH(CH_3)_2, 0.81 (3H, d, J = 6Hz, CH(C_{\underline{H}_3})_2), and 0.74
20
      (3H, d, J = 6Hz, CH(CH<sub>3</sub>)<sub>2</sub>).
21
- 22
      delta<sub>C</sub> (63.9MHz, D<sub>6</sub>-DMSO) 173.0, 171.0, 168.8, 139.8,
23
      138.0, 130.5, 129.0, 128.5, 127.5, 125.8, 125.5, 54.2,
24
      46.0, 45.5, 38.0, 31.5, 25.5, 25.2, 24.7, and 21.0.
25
26
      Example 17
27
28
      [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-chlorophenylthio-
29
```

methyl) succinyl]-L-phenylalanine-N-methylamide

WO 90/05719 PCT/GB89/01399

53

Prepared by the method described in example 1g to give

```
material with the following characteristics.
 2
 3
 4
    m.p. 231-234°C
 5
 6
     [alpha]_D = -96.5^{\circ}
 7
    Analysis calculated for C25H32ClN3O4S
 8
 9
     Requires: C59.34 H6.37 N8.30
10
     Found:
               C59.51 H6.43 N8.24
11
     delta_{H} (250MHz, D_{6}-DMSO) 8.85 (1H, s, N\underline{H}OH), 8.37 (1H,
12
     d, J = 8.5Hz, CONH), 7.90 (1H, m, CONHMe), 7.30 - 6.88
13
     (9H, m, aromatic H), 4.66 (1H, m, CHCH2Ph), 2.96 (1H,
14
15
     bd, J = 14Hz, CHCH_2Ph), 2.76 (1H, bt, J = 13Hz,
16
     CHCH_2Ph) 2.60 (3H, d, J = 5Hz, NHCH_3), 2.55 - 2.40 (2H,
     m, CH_2S), 2.16 (1H, m, CHCO), 2.01 (1H, d, J = 14Hz,
17
     CHCO), 1.37 (2H, m), 0.91 (1H, m, CH_2CH(CH_3)_2), 0.81
18
     (3H, d, J = 6Hz, CH(CH<sub>3</sub>)<sub>2</sub>), and 0.74 (3H, d, J =
19
20.
     6Hz, CH(CH_3)_2).
21
     delta<sub>C</sub> (63.9MHz, D<sub>6</sub>-DMSO) 172.7, 171.6, 168.1, 139.2,
22
23
     138.1, 130.3, 129.2, 127.9, 126.2, 125.9, 125.5, 125.0,
     54.1, 46.3, 45.8, 37.8, 32.0, 25.7, 25.2, 24.2, and
24
25
     21.7.
26
27
28
29
30
31
32
33
```

```
Example 18
```

1 .2 3

[4-(N-Hydroxyamino)-2R-isobuty1-3S-(3-methylphenylthiomethyl) succinyl]-L-phenylalanine-N-methylamide

Prepared by the method described in example 1g to give material with the following characteristics.

18 Analysis calculated for $C_{26}H_{35}N_3O_4S$

19 Requires: C64.30 H7.26 N8.65

20 Found: C63.81 H7.21 N8.48

delta_H (250MHz, D_6 -DMSO) 8.83 (1H, s, NHOH), 8.35 (1H, d, J = 8.5Hz, CONH), 7.86 (1H, m, CONHMe), 7.28 - 6.77(9H, m, aromatic H), 4.66 (1H, m, CHCH₂Ph), 2.96 (1H, dd, J = 14,4Hz, $CHCH_2Ph$), 2.80 (1H, bt, J = 13Hz, $CHCH_{2}Ph)$ 2.59 (3H, d, J = 5Hz, $NHCH_{3}$), 2.55 - 2.37 (2H, m, CH_2S), 2.16 (2H, m, 2xCHCO), 1.38 (2H, m), 0.91 (1H, m, $C_{\underline{H}_2}CH(CH_3)_2$), 0.81 (3H, d, J = 6Hz, $CH(C_{\underline{H}_3})_2$), and 0.74 (3H, d, J = 6Hz, $CH(CH_3)_2$).

WO 90/05719 PCT/GB89/01399

```
Example 19
```

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-acetyl)-aminophenylthiomethyl)succinyl]-L-phenylalanine-N-methylamide.

16 A) [2R-isobutyl-3S-(4-aminophenylthiomethyl)succinyl]-17 L-phenylalanine -N-methylamide.

19 Prepared by the method described in example 1f to give $_{20}$ material with the following characteristics.

```
delta_{H} (250MHz, D_{6}-DMSO) 8.27 (1H, d, J = 8.5Hz, CON_{H}),
22
    7.81 (1H, m, CONHMe), 7.30 - 7.00 (5H, m, phenyl H),
23
    6.86 (2H, d, J = 8.5Hz, aromatic H), 6.45 (2H, d, J =
24
    8.5Hz, aromatic H), 5.25 (1H, bs, CO_{2}H), 4.48 (1H, m,
25
    CHCH_2Ph), 2.91 (1H, dd, J = 14,4Hz, CHCH_2Ph), 2.88 (1H,
26
    dd, J = 14,10Hz, CHCH_2Ph) 2.56 (3H, d, J = 5Hz, NHCH_3),
27
    2.43 - 2.24 (3H, m, C_{H_2}S and C_{H_2}CO), 2.03 (1H, d, J =
28
   10Hz, CHCO), 1.41 (1H, t, J = 11Hz, CH_2CH(CH_3)_2), 1.26
29
    (1H, m, CH_2CH(CH_3)_2), 0.85 (1H, m, CH_2CH(CH_3)_2), 0.81
30
    (3H, d, J = 6Hz, CH(CH<sub>3</sub>)<sub>2</sub>), and 0.74 (3H, d, J=6Hz,
31
    CH(CH_3)_2).
32
```

B) [2R-isobutyl-3S-(4-(N-acetyl)aminophenyl-thiomethyl) - succinyl]-Lphenylalanine-N-methylamide. 3 . The product from above (350mg, 0.74 mmol) was dissolved 4 in DCM (5 ml) cooled in an ice bath then triethylamine 5 (75mg, 0.74 mmol), DMAP (91mg, 7.4 mmol) and finally 6 acetic anhydride (83mg, 8.2 mmol) were added and the 7 solution stirred at RT for 90 minutes. The mixture was 8 partitioned between ethyl acetate and citric acid then 9 the organic layer washed with water and finally dried 10 over magnesium sulphate. Solvent removal gave the crude 11 product as pale yellow crystals (160mg, 0.31 mmol, 12 13 14 delta_H (250MHz, D₆-DMSO) 9.94 (1H, s, CO₂H), 8.34 (1H, 15 d, J = 8.5 Hz, CONH), 7.90 (1H, m, CONHMe), 7.46 (2H, d, 16 J = 8.5Hz, aromatic H) 7.30 - 7.00 (5H, m, phenyl H), 17 6.96 (2H, d, J = 8.5Hz, aromatic H), 4.57 (1H, m, 18 $CHCH_2Ph$), 2.91 (1H, dd, J = 14,4Hz, $CHCH_2Ph$), 2.88 (1H, 19 bt, J = 13Hz, $CHCH_2Ph$), 2.58 (3H, d, J = 5Hz, $NHCH_3$), 20 2.43 - 2.16 (3H, m, CH_2S and CHCO), 2.10 (1H, d, J = 21 14Hz, CHCO), 1.35 (1H, t, J = 14Hz, $CH_2CH(CH_3)_2$), 1.26 22 (1H, m, $CH_2CH(CH_3)_2$), 0.86 (1H, m, $CH_2CH(CH_3)_2$), 0.81 23 (3H, d, J = 6Hz, CH(CH_3)2), and 0.74 (3H, d, J = 24 6Hz, $CH(CH_3)_2$). 25 26 [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-acetyl)-27 aminophenylthiomethyl)succinyl]-L-phenylalanine-N-28 methylamide. 29 30 Prepared by the method described in example 1g to give 31 material with the following characteristics. 32 33

PCT/GB89/01399

m.p. 201 -202°C (dec.) 1 2 $[alpha]_D = -7.5^{\circ}$ (c=1.0, methanol) 3 4 delta_H (250MHz, D₆-DMSO) 9.90 (1H, s, NHOH), 8.82 (1H, 5 s, NHOH), 8.30 (1H, d, J = 8.5Hz, CONH), 7.85 (1H, m, 6 CONHMe), 7.45 (2H, d, J = 8.5Hz, aromatic H), 7.28 -7 6.94 (5H, m, phenyl H), 6.90 (2H, d, J = 8.5Hz, 8 aromatic H), 4.66 (1H, m, $CHCH_2Ph$), 2.90 (1H, dd, J =14,4Hz, CHCH₂Ph), 2.76 (1H, bt, J = 13Hz, CHCH₂Ph), 10 2.50 (3H, d, J = 5Hz, $NHCH_3$), 2.49 - 2.35 (2H, m, 11 CH_2S), 2.14 (1H, m, CHCO), 2.03 (4H, s + m, $COCH_3$ and 12 CHCO), 1.35 (2H, m), 0.86 (1H, m, $CH_2CH(CH_3)_2$), 0.81 13 (3H, d, J = 6Hz, CH(CH₃)₂), and 0.74 (3H, d, J = 6Hz,14 $CH(CH_3)_2).$ 15

16 17

Example 20

18 19

[4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulfinyl-methylsuccinyl]-L-phenylalanine-N-methylamide.

20 21 22

2324252627

28 29

30

31 [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylthiomethyl-32 succinyl]-L-phenylalanine-N-methylamide (250mg, 33 0.53mmol) was dissolved in methanol (50 ml) and metachloroperbenzoic acid (100mg, 0.58 mmol) was added.

```
After stirring for 1h at room temperature ether was
 2
    added and the mixture filtered.
                                        Solvent removal gave
 3
    the crude white solid which was recrystallised from
    methanol / water then slurried in ether to remove final
 5
    traces of meta-chlorobenzoic acid to give the desired
   material (70 mg, 0.014 mmol, 27%).
 8
    m.p. 186 -188<sup>0</sup>C
 9
10
    [alpha]_D = -13.6^{\circ} (c=0.5, methanol)
11
12
    Analysis calculated for C25H33N3O5S.0.5H2O
13
14
    Requires: C60.46 H6.90 N8.46
               C60.58 H6.69 N8.29
    Found:
15
16
    delta<sub>H</sub> (250MHz, D<sub>6</sub>-DMSO, mixture of diastereomers) 9.04
17
    + 8.93 (1H, 2xs, NHOH), 8.29 + 8.16 (1H, 2xd, J = 8.5
18
    Hz, CONH), 7.79 (1H, m, CONHMe), 7.90 - 7.40 (8H, m,
19
    aromatic H), 7.06 + 6.82 (2H, 2xm, SO-Aromatic), 4.37
20
    (1H, m, CHCH<sub>2</sub>Ph), 2.93 - 2.58 (3H, m, containing
21
    CHCH_2Ph), 2.52 (3H, m, NHCH_3), 2.49 + 2.37 (1H, 2xm),
22
    1.49 - 1.25 (2H, m, CH_2CH(CH_3)_2 and CH2CH(CH_3)_2), 0.95
23
24
    (1H, m, CH_2CH(CH_3)_2), 0.81 (3H, d, J = 6Hz, CH(CH_3)_2),
    and 0.74 (3H, d, J=6Hz, CH(CH_3)_2).
25
26
             (63.9MHz, D<sub>6</sub>-DMSO, mixture of diastereomers)
    delta
27
    172.2, 171.4, 171.3, 167.7, 144.5, 138.0, 137.9, 131.3,
28
    130.9, 129.6, 129.3, 129.1, 128.8, 128.3, 127.8, 126.5,
29
    126.2, 124.3, 123.6, 59.8, 58.1, 54.3, 54.0, 46.2,
30
    45.8, 41.6, 40.9, 37.6, 37.4, 25.6, 25.0, 24.3, 24.2,
31
    21.7, and 21.6.
32
33
```

32

33

```
Example 21
 1
 2
    [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulfonyl-
 3
    methylsuccinyl]-L-phenylalanine-N-methylamide.
 4
 5
 6
 7
 8
 9
10
11
12
    [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylthiomethyl-
13
    succinyl]-L-phenylalanine-N-methylamide (50mg,
14
    0.11mmol) was dissolved in methanol (12 ml) and meta-
15
    chloroperbenzoic acid (40mg, 0.23 mmol) was added.
16
    After stirring for 3h at room temperature ether was
17
    added and the mixture filtered.
                                        Solvent removal gave
18
    the crude white solid which was slurried in ether to
19
    remove final traces of meta-chlorobenzoic acid to give
20
    the desired material.
21
22
    m.p. 228 - 231^{\circ}C
23
24
    [alpha]_D = 16.8^{\circ} (c=0.5, methanol)
25
26
    Analysis calculated for C_{25}H_{33}N_3O_6S.0.3H_2O
27
    Requires: C58.99 H6.65 N8.25
28
              C58.92 H6.51 N8.05
29
   Found:
30
   delta<sub>H</sub> (250MHz, D<sub>6</sub>-DMSO) 8.66 (1H, s, NHOH), 8.25 (1H,
```

d, J = 8.5 Hz, CONH), 7.83 (1H, m, CONHMe), 7.75 - 7.50

(5H, m, aromatic H), 7.30 7.05 (5H, m, aromatic H),

4.36 (1H, m, $CHCH_2Ph$), 2.86 (1H, dd, J = 14.5 Hz, $CHCH_2Ph)$, 2.75 (1H, dd, J = 14,10 Hz, $CHCH_2Ph$), 2.54 (3H, d, J = 4.5 Hz, NHCH₃), 2.54 (2H, m), 1.30 (2H, m,3 $CH_2CH(CH_3)_2$ and $CH_2CH(CH_3)_2)$, 0.86 (1H, m, $CH_2CH(CH_3)_2$, 0.75 (3H, d, J = 6Hz, $CH(CH_3)_2$), and 0.71 (3H, d, J = 6Hz, CH(CH₃)₂).6 7 Example 22 8 9 10 [4-(N-Hydroxyamino)-2R-isobuty1-3Sthiophenylsulphinylmethyl-succinyl] -L-phenylalanine-N-11 methylamide 12 13. 14 15 16 17 СОИНОН 18 19 20 21. [4-(N-Hydroxyamino)-2R-isobutyl-3S-thiophenylthio-22 methyl-succinyl]-L-phenylalanine-N-methylamide (50mg, 23 0.11mmol) was treated as described in example 21 to 24 25. yield the title compound (16mg, 0.03 mmol, 29%) as a 26 mixture of diastereomer with the following 27 characteristics: 28 m.p. 195 -197°C (dec.) 29 30

Analysis calculated for $C_{23}H_{31}N_3O_5S_2.0.5H_2O$ 31

Requires: C54.96 H6.42 N8.36 32

Found: 33 C54.91 H6.23 N8.23

PCT/GB89/01399

WO 90/05719

delta_H (250MHz, D₆-DMSO, mixture of diastereomers) 9.04 + 8.96 (1H, 2xs, NHOH), 8.34 + 8.29 (1H, 2xd, J = 8.52 Hz, CONH), 8.02 + 7.98 (1H, 2xm, CONHMe), 7.81 (1H, bs, thiophene-H), 7.42 (1H, s, thiophene-H), 7.25 - 7.15 4 (5H, m, phenyl), 7.03 (1H, bs, thiophene-H), 4.43 (1H, 5 m, $CHCH_2Ph$), 3.0 - 2.6 (4H, m, containing $CHCH_2Ph$), 6 2.52 (7H, m, containing NHC \underline{H}_3), 2.05 (1H, m), 1.6 - 1.2 7 (2H, m, $CH_2CH(CH_3)_2$ and $CH_2CH(CH_3)_2$), 0.87 (1H, m, 8 $CH_2CH(CH_3)_2$, and 0.85 - 0.71 (6H, m, $CH(CH_3)_2$). 9

61

10

Example 23

11 12

[4-(N-Hydroxyamino)-2R-isobutyl-3S-13 thiophenylsulphonylmethyl-succinyl] -L-phenylalanine-N-14 methylamide. 15

16 17

23 24

25

26

27

28

[4-(N-Hydroxyamino)-2R-isobutyl-3S-thiophenylthiomethyl-succinyl]-L-phenylalanine-N-methylamide (75mg, 0.16mmol) was treated as described in example 22 to yield the title compound (40mg, 0.08 mmol, 49%) with the following characteristics:

29 30

32

Analysis calculated for C23H31N3O6S2 33

```
Requires: C54.21 H6.13 N8.24
    Found:
               C54.07 H6.19 N8.04
 3
    delta_{H} (250MHz, D_{6}-DMSO) 887 (1H, s, N_{H}OH), 8.25 (1H,
    d, J = 8.5 \text{ Hz}, CONH), 8.09 (1H, d, J = 4.7 \text{ Hz},
 5
   thiophene-H), 7.83 (1H, m, CONHMe), 7.53 (1H, d, J = 3)
    Hz, thiophene H), 7.25 - 7.12 (6H, m, phenyl and
    thiophene-H), 4.36 (1H, m, CHCH<sub>2</sub>Ph), 3.38 (1H, dd, J =
    14,11 Hz, SCH_2), 2.87 (1H, dd, J = 14,5 Hz, CHCH_2Ph),
10 2.75 (1H, dd, J = 14,10 Hz, CHCH_2Ph), 2.70 - 2.36 (6H,
    m, containing NHC\underline{H}_3), 1.20 (2H, m, \underline{CH}_2CH(CH_3)_2 and
11
    CH_2CH(CH_3)_2), 0.89 (1H,m, CH_2CH(CH_3)_2), and 0.75 (6H,
12
    m, CH(CH_3)_2).
13
14
    delta<sub>C</sub> (63.9MHz, D<sub>6</sub>-DMSO) 172.0, 171.2, 166.5, 140.0,
15
    138.0, 135.4, 134.6, 129.0, 128.4, 128.2, 126.6, 54.3,
16
    45.6, 37.5, 25.6, 25.0, 24.2, and 21.7.
17
18
    Example 24
19
20
    [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulfonyl-
21
    methylsuccinyl]-L-phenylalanine-N-methylamide sodium
22
    salt.
23
24
25
26
27
                              CONHONa
28
29
```

30 31

[4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulfonyl-

methylsuccinyl]-L-phenylalanine-N-methylamide (50mg, 0.1mmol) was dissolved in methanol (10ml) and sodium hydroxide solution (0.1M, 1.0ml) added to give a homogeneous solution. The methanol was removed under reduced pressure then the residual aqueous solution freeze dried to give the title compound (40mg).

7

delta_H (250MHz, D₆-DMSO) 8.66 (1H, s, NHOH), 8.25 (1H, 8 d, J = 8.5 Hz, CONH), 7.83 (1H, m, CONHMe), 7.75 - 7.50 9 (5H, m, aromatic H), 7.30 7.05 (5H, m, aromatic H), 10 4.36 (1H, m, $CHCH_2Ph$), 2.86 (1H, dd, J = 14,5 Hz, 11 $CHCH_2Ph$), 2.75 (1H, dd, J = 14,10 Hz, $CHCH_2Ph$), 2.54 12 (3H, d, J=4.5 Hz, $NHCH_3$), 2.54 (2H, m), 1.30 (2H, m, 13 $CH_2CH(CH_3)_2$ and $CH_2CH(CH_3)_2)_1$ 0.86 (1H, 14 $C_{\underline{H}_2}CH(CH_3)_2$, 0.75 (3H, d, J = 6Hz, $CH(C_{\underline{H}_3})_2$), and 0.71 15 (3H, d, J = 6Hz, CH(CH₃)₂).16

17 18

Example 25

19 20

21

22

[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(isobutyloxy-carbonylamino)phenyl)thiomethyl-succinyl]-L-phenyl-alanine-N-methylamide

2324252627282930

31 32

33 a) [4-Hydroxy-2R-isobutyl-3S-(4-aminophenyl)thio-

```
1 methylsuccinyl]-L-phenylalanine-N-methylamide was
    prepared by the method described in example 1f to give
    a compound with the following characteristics.
 3
   delta_{H} (250MHz, D_{6}-DMSO) 8.26 (1H, d, J = 8.5 Hz,
    CONH), 7.81 (1H, m, CONHMe), 7.27 - 7.15 (5H, m, phenyl
    H), 6.85 (2H, d, J = 8.5Hz, aromatic H), 6.46 (2H, d, J
 8 = 8.5Hz, aromatic H), 5.2 (1H, bs, CO_{2}H), 4.48 (1H, m,
    CHCH_2Ph), 2.90 (1H, dd, J = 13.5,4.3 Hz, CHCH_2Ph), 2.75
    (1H, dd, J = 13.6, 10 Hz, CHCH_2Ph), 2.56 (3H, d, J =
11 4.5 Hz, NHCH3), 2.50 - 2.25 (3H, m), 2.03 (1H, d, J =
    10 Hz), 1.41 (1H, m, CH_2CH(CH_3)_2), 1.26 (1H, m,
12
   CH_2CH(CH_3)_2), 0.86 (1H, m, CH_2CH(CH_3)_2), 0.75 (3H, d, J
    = 6Hz, CH(CH_3)_2, and 0.71 (3H, d, J = 6Hz, CH(CH_3)_2).
14
15
16 b) N,N-Dimethylglycine (100mg, 0.97 mmol) was stirred
    in dry THF (50ml) and triethylamine (108mg, 1.1mmol)
    and isobutylchloroformate (146mg, 1.1mmol) were added.
18
    After 1h the product from example 26a (500mg, 1.1mmol)
19
   was addedand the mixture stirred for a further 1h. The
20
    reaction was worked up by partitioning between citric
21
   acid and ethyl acetate, drying the organic layer and
22
   solvent removal to give the crude product (1g).
23
   Solution of the crude solid in ethyl acetate then
24
   precipitation with ether resulted in white crystals of
25
   the isobutylchloroformate derivative.
26
27
        [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(isobutyloxy-
28
   carbonylamino) phenyl)thiomethyl-succinyl]-L-phenyl-
29
   alanine-N-methylamide
30
```

The product from example 26b was converted to the 32

hydroxamic acid as described in example 1g. to give a 33 compound with the following characteristics.

```
m.p. 198 - 200^{\circ}C
   [alpha]_D = -8.5^{\circ} (c=1, methanol)
    Analysis calculated for C30H42N4O6S
    Requires: C61.41 H7.22 N9.55
              C62.04 H7.32 N9.67
    Found:
 7
 8
    delta_{H} (250MHz, D_{6}-DMSO) 9.60 (1H, s, NHOH), 8.83 (1H,
 9
    s, NHOH), 8.31 (1H, d, J = 8.5 Hz, CONH), 7.85 (1H, m,
10
    CONHMe), 7.36 - 7.25 (4H, m, aromatic H), 7.14 - 7.05
11
    (3H, m, aromatic H), 6.91 (2H, d, J = 8.5Hz, aromatic
12
    H), 4.56 (1H, m, CHCH_2Ph), 3.87 (2H, d, J = 7Hz,
13
    OCH_2CH(CH_3)_2), 2.92 (1H, dd, J = 13.7,4.0 Hz, CHCH_2Ph),
14
    2.76 (1H, dd, J = 13.6,10 \text{ Hz}, CHCH_2Ph), 2.58 (3H, d, J
15
    = 4.5 \text{ Hz}, NHCH_3), 2.50 - 2.34 (2H, m), 2.16 - 1.87 (3H,
16
    m), 1.35 (2H, m, C\underline{H}_2CH(CH_3)_2 and CH_2C\underline{H}(CH_3)_2), 0.93
17
           d, J = 6.6Hz, OCH_2CH(CH_3)_2, 0.87 (1H,m,
18
    CH_2CH(CH_3)_2), 0.75 (3H, d, J = 6Hz, CH(CH_3)_2), and
19
    0.71 (3H, d, J = 6Hz, CH(CH_3)_2).
20
21
22
    Example 26
23
24
    [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-methyl-N-
25
    (tertbutoxycarbonyl)-glycylamino) phenyl)thiomethyl-
26
    succinyl]-Lphenylalanine-N-methylamide.
27
28
29
30
```

```
[4-Hydroxy-2R-isobutyl-3s-(4-(N-methyl-N-(tert-
    butoxycarbonyl)glycylamino) phenyl)thiomethyl-
 2
    succinyl]-L-phenylalanine-N-methylamide was prepared as
 3
    described in example 26b by substitution of N-BOC
 4
    sarcosine for the acid component.
 5
 6
    delta<sub>H</sub> (250MHz, D_6-DMSO) 9.97 (1H, s, CO_2H), 8.36 (1H,
 7
    d, J = 8.5 \text{ Hz}, CONH), 7.91 (1H, m, CONHMe), 7.48 (2H,
 8
    d, J = 8.5Hz, aromatic H), 7.40 - 7.05 (5H, m, aromatic
 9
    H), 6.97 (2H, d, J = 8.5Hz, aromatic H), 4.58 (1H, m,
10
    CHCH_2Ph), 3.95 (2H, d, J = 9Hz, NCH_2CO), 2.92 (4H, m+d,
11
    CHCH_2Ph and BOCNCH_3), 2.76 (1H, dd, J = 13,10 Hz,
12
    CHCH_2Ph), 2.58 (3H, d, J = 4.5 Hz, NHCH_2), 2.50 - 2.09
13
    (4H, m),
               1.46 - 1.33 (11H,
                                      m + 2xs
14
    CH_2CH(CH_3)_2 and CH_2CH(CH_3)_2),
                                           0.87
                                                   (1H,
15
    CH_2CH(CH_3)_2), 0.75 (3H, d, J = 6Hz, CH(CH_3)_2), and
16
    0.71 (3H, d, J = 6Hz, CH(CH_3)_2).
17
18
        [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-methyl- N-
    b)
19
    (tertbutoxycarbonyl)-glycylamino)phenyl)- thiomethyl-
20
    succinyl]-Lphenylalanine-N-methylamide was prepared
21
    from the material produced in example 27a as described
22
    in example 1g.
23
24
    delta_{H} (250MHz, D_{6}-DMSO) 9.97 (1H, s, CONHOH), 8.83
25
    (1H, s, NHOH), 8.32 (1H, d, J = 8.5 Hz, CONH), 7.86
26
    (1H, m, CONHMe), 7.46 (2H, d, J = 8.5Hz, aromatic H),
27
    7.28 - 7.00 (5H, m, aromatic H), 6.97 (2H, d, J =
28
    8.5Hz, aromatic H), 4.56 (1H, m, CHCH<sub>2</sub>Ph), 3.94 (2H, d,
29
    J = 9Hz, NCH_2CO), 2.87 (4H, m+d, CHCH_2Ph and BOCNCH_3),
30
    2.76 (1H, m, CHCH<sub>2</sub>Ph), 2.57 (3H, d, J = 4.5 Hz, NHCH<sub>3</sub>),
31
    2.25 - 1.91 (2H, m), 1.42 - 1.30 (11H, m + 2xs,
32
              CH_2CH(CH_3)_2 and CH_2CH(CH_3)_2, 0.92 (1H, m,
    (CH_3)_3C,
33
   CH_2CH(CH_3)_2), 0.80 (3H, d, J = 6Hz, CH(CH_3)_2), and
```

0.73 (3H, d, J=6Hz, $CH(CH_3)_2$).

PCT/GB89/01399 WO 90/05719

67

1 2 Example 27 3 4 Collagenase inhibition activity 5 The potency of compounds of general formula I to act 6 7 as inhibitors of collagenase (a metalloproteas involved in tissue degradation) was determined by the 8 9 procedure of Cawston and Barrett, (Anal. Biochem., 99, 340-345, 1979), hereby incorporated by reference, 10 whereby a 1mM solution of the inhibitor being tested or 11 dilutions thereof was incubated at 370 for 16 hours 12 with collagen and collagenase (buffered with 25mM 13 Hepes, pH 7.5 containing 5mM CaCl2, 0.05% Brij 35 and 14 0.02% NaNa). The collagen was acetylated 14C collagen 15 prepared by the method of Cawston and Murphy 16 in Enzymology, 80, 711, 1981), hereby incorporated by 17 reference. The samples were centrifuged to sediment 18 undigested collagen and an aliquot of the radioactive 19 supernatant removed for assay on a scintillation 20 counter as a measure of hydrolysis. The collagenase 21 activity in the presence of 1 mM inhibitor, or a 22 23 dilution thereof, was compared to activity in a control devoid of inhibitor and the results reported below as 24 that inhibitor concentration effecting 50% inhibition 25 of the collagenase (IC_{50}). 26 27

28	Compound of Example No.	<u>IC</u> 50
29	1	20 nM
30	2	8 nM
31	5 6	.3 nM (50% @ 1 mcM)
32		, 230 G T MCPT

inhibitors of stromelysin was determined using the 7 procedure of Cawston et al (Biochem. J., 195, 159-165 1981), hereby incorporated by reference, whereby a 1mM solution of the inhibitor being tested or dilutions thereof was incubated at 37°C for 16 hours with stromelysin and ^{14}C acetylate casein (buffered with 25mM Hepes, pH 7.5 containing 5mM CaCl₂, 0.05% Brij 35 13 and 0.02% NaN_3 . The casein was ^{14}C acetylated 14 according to the method described in Cawston et al 15 (Biochem. J., 195, 159-165, 1981), hereby incorporated 16 by reference. The stromelysin activity in the presence 17 of 1mM, or a dilution thereof, was composed to activity 18 in a control devoid of inhibitor and the results 19 reported below as that inhibitor concentration 20 effecting 50% inhibition of the stromelysin (IC_{50}). 21

22

26 27

Examples of unit dosage compositions are as follows:

28 29

30

31

32

33 -

WO 90/05719 PCT/GB89/01399

```
1
 2
 3
    Example 29
 5
         Capsules:
 6
                                             Per 10,000
 7
           Ingredients
                              Per Capsule
                                             Capsules
 8
 9
              Active ingredient
         1.
10
              Cpd. of Form. I
                                40.0 mg
                                                  400 g
11
              Lactose
                                 150.0 mg
                                                 1500 g
         2.
12
              Magnesium
         3.
13
              stearate
                                  4.0 mg
14
                                 194.0 mg
                                                 1940 g
15
16
    Procedure for capsules:
17
18
              Blend ingredients No. 1 and No. 2 in a
    Step 1.
19
              suitable blender.
20
              Pass blend from Step 1 through a No. 30 mesh
21
    Step 2.
              (0.59 mm) screen.
22
              Place screened blend from Step 2 in a
    Step 3.
23
              suitable blender with ingredient No. 3 and
24
              blend until the mixture is lubricated.
25
              Fill into No. 1 hard gelatin capsule shells
    Step 4.
26
              on a capsule machine.
27
28
29
30
31
32
33
```

1	Example 30		
2		•	
3	Tablets:		
4		Per 10,000	
5		<u>Ingredients</u> <u>Per Tablet</u> <u>Tablets</u>	
6	•		
7	1.	Active ingredient	
8	•	Cpd. of Form. I 40.0 mg 400 g	
9 .	2.	Corn Starch 20.0 mg 200 g	
10	3.	Alginic acid 20.0 mg 200 g	
11	. 4.	Sodium alginate 20.0 mg 200 g	
12	5.	Magnesium	
13		stearate 1.3 mg 13 g	
14		101.3 mg 1013 g	
15			
16	Procedure	for tablets:	
17	Step 1.	Blend ingredients No. 1, No. 2, No. 3 and No.	
18		4 in a suitable mixer/blender.	
19	Step 2.	Add sufficient water portionwise to the blend	
20	•	from Step 1 with careful mixing after each	
21		addition. Such additions of water and mixing	
22		until the mass is of a consistency to permit	
23		its conversion to wet granules.	
24	Step 3.	The wet mass is converted to granules by	
25		passing it through an oscillating granulator	
26	·. ·	using a No. 8 mesh (2.38) screen.	
27	Step 4.	The wet granules are then dried in an oven at	
28		140 ^O F (60 ^O C) until dry.	
29	Step 5.	The dry granules are lubricated with	
30	٠.	ingredient No. 5.	
31	Step 6.	The lubricated granules are compressed on a	
32		suitable tablet press.	
33			

WO 90/05719 PCT/GB89/01399

1	Example 3	<u>1</u>		
2				
3	Intr	amuscular Injection:		
4		<u>Ingredient</u>	Per ml.	<u>Per liter</u>
5	1.	Compound of Formula	I	
6		Active ingredient	10.0 mg	10 g
7	2.	Istonic buffer		
8		solution pH 4.0.	q.s.	q.s.
9				
10	Procedure			
11	Step 1.	Dissolve the active	ingredient i	n the buffer
12		solution.	1	
13	Step 2.	Aseptically filter		
14	Step 3.	The sterile solution		tically
15		filled into sterile		
16	Step 4.	The ampoules are sea	aled under as	petic
17		conditions.		
18				
19	Example 3			
20				
21	Supp	oositories:		
22				Per
23		<u>Ingredients</u>	Per Supp.	1,000 Supp
24	1.	Compound of Form. I		
25		Active ingredient	40.0 mg	40 g
26	2.	Polyethylene Glycol	•	
27		1000	1350.0 mg	1,350 g
28	3.	Polyethylene Glycol		
29		4000	450.0 mg	<u>450 q</u>
30			1840.0 mg	1,840 g
31				
32			•	
33				

Procedure:
Step 1. Melt ingredient No. 2 and No. 3 together and
stir until uniform.
Step 2. Dissolve ingredient No. 1 in the molten mass
from Step 1 and stir until uniform.
Step 3. Pour the molten mass from Step 2 into
suppository moulds and chill.
Step 4. Remove the suppositories from moulds and
wrap.
Example 33
Eye Ointment
An appropriate amount of a compound of general formula
I is formulated into an eye ointment base having the
following composition:
Liquid paraffin 10%
Wool fat 10%
Yellow soft paraffin 80%
Example 34
Topical skin ointment
An appropriate amount of a compound of general formula
I is formulated into a topical skin ointment base
having the following composition:
Emulsifying wax 30%
White soft paraffin 50%
Liquid paraffin 20%

(I)

<u>CLAIMS</u>

1 2 3

1. A compound of general formula I:

4 5

6 7 8

9

 R^2 N H R^3 R^4 R^5 R^5

10
11 wherein:

12

13 R^1 represents a C_1 - C_6 alkyl, phenyl, thiophenyl, 14 substituted phenyl, phenyl(C_1 - C_6) alkyl, 15 heterocyclyl, (C_1 - C_6) alkylcarbonyl or phenacyl or 16 substituted phenacyl group; or when n = 0, R^1 17 represents SR^X , wherein R^X represents a group:

18

23 24

25

represents a hydrogen atom or a C_1 - C_6 alkyl, C_1 - C_6 alk enyl, phenyl(C_1 - C_6) alkyl, cycloalkyl(C_1 - C_6) alkyl or cycloalkenyl(C_1 - C_6) alkyl group;

30

represents an amino acid side chain or a C_1-C_6 alkyl, benzyl, (C_1-C_6) alkoxy) benzyl or benzyloxy (C_1-C_6) alkyl) or benzyloxy benzyl group;

12

16

- R^4 represents a hydrogen atom or a C1-C6 alkyl group; 2 R^5 3 represents a hydrogen atom or a methyl group; is an integer having the value 0, 1 or 2; and 5 n 6 7 represents a C_1 - C_6 hydrocarbon chain, optionaly substituted with one or more c_1-c_6 alkyl, phenyl 8 9 or substituted phenyl groups; 10 11 or a salt thereof.
- 13 2. A compound as claimed in Claim 1, in which the 14 chiral centre adjacent the substituent R^3 has S 15 stereochemistry.
- 17 3. A compound as claimed in Claim 1 or 2, wherein the 18 chiral centre adjacent the substituent \mathbb{R}^2 has \mathbb{R} 19 stereochemistry.
- 4. A compound as claimed in Claim 1, 2 or 3, in which R¹ represents a hydrogen atom or a C₁-C₄ alkyl, phenyl, thiophenyl, benzyl, acetyl or phenacyl group.
- 25 5. A compound as claimed in any one of Claims 1 to 4, wherein ${\bf R}^2$ represents a ${\bf C}_3{\bf -C}_6$ alkyl group.
- 28 6. A compound as claimed in any one of Claims 1 to 5, 29 wherein \mathbb{R}^3 represents a benzyl or 30 $4-(C_1-C_6)$ alkoxyphenylmethyl or benzyloxybenzyl group. 31
- 7. A compound as claimed in any one of Claims 1 to 6, wherein \mathbb{R}^4 represents a C_1-C_4 alkyl group.

WO 90/05719 PCT/GB89/01399

```
A compound as claimed in any one of Claims 1 to 7,
 1
     wherein R<sup>5</sup> represents a hydrogen atom.
 2
 3
          [4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenylthio-
     9.
 4
     methyl)-succinyl]-L-phenylalanine-N-methylamide,
 5
 6
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiophenylthio-
 7
     methyl) succinyl]-L-phenylalanine-N-methylamide,
 8
 9
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(benzylthiomethyl)
10
     succinyl]-L-phenylalanine-N-methylamide,
11
12
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(acetylthiomethyl)
13
     succinyl]-L-phenylalanine-N-methylamide or
14
15
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiolmethyl)
16
     succinyl]-L-phenylalanine-N-methylamide
17
18
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(pivaloylthiomethyl)
19
     succinyl]-L-phenylalanine-N-methylamide
20
21
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(phenylthiomethyl)
22
     succinyl]-L-phenylalanine-N-methylamide sodium salt
23
24
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-methoxyphenyl-
25
     thiomethyl) succinyl]-L-phenylalanine-N-methylamide
26
27
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-hydroxyphenyl-
28
     thiomethyl)succinyl]-L-phenylalanine-N-methylamide
29
3.0
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(2-thiophenethio-
31
     methyl)succinyl]-L-phenylalanine-N-methylamide sodium
32
33
     salt
```

```
[4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-methoxyphenyl-
    thiomethyl) succinyl]-L-phenylalanine-N-methylamide
 3
     sodium salt
 4
 5
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-tertbutylphenyl-
 6
     thiomethyl) succinyl]-L-phenylalanine-N-methylamide
 7
 8
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(2,4-dimethylphenyl-
 9.
     thiomethyl) succinyl]-L-phenylalanine-N-methylamide
10
11
     bis-S,S'-{ [4 (N-Hydroxyamino-2R-isobutyl-3S-(thiomethyl)
     succinyl]-L-phenylalanine-N-methylamide disulphide
12
13
14
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-bromophenylthio-
     methyl) succinyl]-L-phenylalanine-N-methylamide
15
16
17 . .
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-chlorophenylthio-
18
     methyl) succinyl]-L-phenylalanine-N-methylamide
19
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(3-methylphenylthio-
20
21
     methyl) succinyl]-L-phenylalanine-N-methylamide
22
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-acetyl)-amino-
23
     phenylthiomethyl) succinyl]-L-phenylalanine-N-methyl-
24
     amide
25
26
27
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulphinyl-
28
     methylsuccinyl]-L-phenylalanine-N-methylamide
29
30
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulphonyl-
31 -
     methylsuccinyl]-L-phenylalanine-N-methylamide
32 .
```

WO 90/05719 PCT/GB89/01399

```
1
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-thiophenylsulphinyl-
     methyl-succinyl]-L-phenylalanine-N-methylamide
 2
 3
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-thiophenylsulphonyl-
 4
     methyl-succinyl]-L-phenylalanine-N-methylamide
 5
 6
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-phenylsulphonyl-
 7
     methyl-succinyl]-L-phenylalanine-N-methylamide sodium
 8
     salt
 9
10
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(isobutyloxy-
11
12
     carbonylamino)phenyl)thiomethyl-succinyl]-L-phenyl-
     alanine-N-methylamide
13
14
15
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(4-(N-methyl-N-
     (tert-butoxycarbonyl)-glycylamino)phenyl)thiomethyl-
16
     succinyl]-L-phenylalanine-N-methylamide
17
18
     or, where appropriate, a salt of such a compound.
19
20
          [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiophenyl-
21
     thiomethyl) succinyl]-L-phenylalanine-N-methylamide, or
22
23
     [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiolmethyl)
24
     succinyl]-L-phenylalanine-N-methylamide
25
26
     or a salt thereof.
27
28
          [4-(N-Hydroxyamino)-2R-isobutyl-3S-(thiophenyl-
29
     thiomethyl)succinyl]-L-phenylalanine-N-methylamide or a
30
31
     salt thereof.
32
33
```

1 12. A compound as claimed in any one of claims 1 to 11

2 for use in human or veterinary medicine.

3

The use of a compound as claimed in any one of

claims 1 to 11 in the preparation of an agent for use 5

in the management of disease involving tissue

degradation and/or in the promotion of wound healing. 7

8

A pharmaceutical or veterinary formulation 9

10 comprising a compound as claimed in any one of claims 1

11 to 11 and a pharmaceutically and/or veterinarily

. 12 acceptable carrier.

13

14 15. A process for preparing a compound of general

15 formula I as defined in claim 1, the process

16 comprising:

17

deprotecting a compound of general formula II 18

 \mathbb{R}^3

19 20

22

23

24 wherein:

25 26

27

 \mathbb{R}^1 , \mathbb{R}^2 , \mathbb{R}^3 , \mathbb{R}^4 , \mathbb{R}^5 , A and n are as defined in

CONHZ

general formula I and Bn represents a

28 benzyloxycarbonyl group; or

29

30 (b) reacting a compound of general formula III

31 32

33

(III)

(II)

PCT/GB89/01399

(III)

wherein:

 R^1 , R^2 , R^3 , R^4 , R^5 , A and n are as defined in general formula I,

with hydroxylamine or a salt thereof; and

optionally after step (a) or step (b) converting a compound of general formula I into another compound of general formula I.

16. A compound of general formula II

(II)

wherein:

 R^1 , R^2 , R^3 , R^4 , R^5 , A and n are as defined in general formula I and Z represents a protecting group.

17. A compound of general formula III

wherein:

 R^1 , R^2 , R^3 , R^4 , R^5 , A and n are as defined in general formula I.

INTERNATIONAL SEARCH REPORT

International Application No

PCT/GB 89/01399

I. CLASSIFICATION OF SUBJECT MATTER (:: several class	ssification sympols apply indicate all) 4		
According to International Patent Classification (IPC) or to both N	07 D 333/34, C 07 C	327/32	
IPC : 317/50, 313/48, A 61 K 31	/13, 31/38		
II. FIELDS SEARCHED			
	nentation Searched ?		
Classification System	Classification Symbols	· · · · · · · · · · · · · · · · · · ·	
IPC ⁵ C 07 C 259/00, 323 C 07 C 327/00, 317	/00, C 07 D 333/00, /00, 313/00		
	r than Minimum Documentation its are included in the Fields Searched ⁹		
	·		
III. DOCUMENTS CONSIDERED TO BE RELEVANT			
Category * Citation of Document, 11 with Indication, where as	ppropriate, of the relevant passages 12	! Relevant to Claim No. 13	
A EP, A, 0236872 (F. HO) 16 September 1987 see claim 1		: 1-17 :	
cited in the applicat:	ion	· .	
A EP, A, 0012401 (MERCK 25 June 1980 see claim 1	& CO. INC.)	: 1-17 :	
cited in the applicati	ion -	•	
A DE, A, 2720996 (E.R. S 24 November 1977 see claim 1 cited in the applicati & US, A, 4105789		1-17	
A EP, A, 0274453 (LABORA 13 July 1988 see claim 1	ATOIRE ROGER BELLON)	1-17	
A EP, A, 0214639 (G.D. S 18 March 1987 see claim 1	SEARLE)	1-17	
*T" later document published after the international filing date or priority date and not in conflict with the application but considered to be of particular relevance. "E" earlier document but published on or after the international filing date. "L" document which may throw doubts on priority claim(s) or which is cried to establish the publication date of another citation or orfer special reason (as specified). "O" document referring to an oral disclosure, use, exhibition or other means. "P" document published after the international filing date but later than the priority date claimed. "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the cannot be considered novel or cannot be considered to involve an inventive step. "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive at particular relevance; the claimed invention cannot be considered to involve an inventive at particular relevance; the claimed invention cannot be considered to involve an inventive at particular relevance; the claimed invention cannot be considered to involve an inventive at particular relevance; the claimed invention cannot be considered novel or cannot be considered novel or cannot be considered novel or cannot be considered to involve an inventive at particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive at particular relevance; the claimed invention or cannot be considered novel or cannot be considered no			
Date of the Actual Completion of the International Search	4		
8th March 1990	1 7 AVR 199	0	
International Searching Authority	Signature of Authorized Officer		
EUROPEAN PATENT OFFICE	AND TO T	Δ7F1 Δ Δ D	

Category *	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No
	cited in the application & US, A, 4599361	:
A	Chemical Abstracts, volume 83, no. 7, 18 August 1975, (Columbus, Ohio, US), J.P. Devlin et al.: "Antibiotic actinonin. III. Synthesis of structural analogs of actinonin by the anhydride-imide method",	1-17
	see page 549, abstract 59249e, & J. Chem. Soc., Perkin Trans. I, 1975, (9), 830-41	i :
		•
:		: !
٠		
		· ·
		,
.•		
:···		

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 8901399

SA 33118

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 04/04/90

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

cited in se	document arch report	Publication date		nt family nber(s)	Publication date
EP-A- (0236872	16-09-87	AU-B- AU-A- JP-A-	588437 6990287 62230757	14-09-89 17-09-87 09-10-87
EP-A- (0012401	25-06-80	AT-T- AU-B- AU-A- CA-C- JP-A- US-A-	E6503 530380 5346179 1262684 55081845 4374829	15-03-84 14-07-83 19-06-80 07-11-89 20-06-80 22-02-83
DE-A- ;	2720996	24-11-77	US-A- CA-A- FR-A, B GB-A- JP-A- US-A- US-A- US-A- US-A- US-A- US-A- US-A- US-A- US-A-	4105789 1103259 2421874 1575850 52136121 4146639 4228184 4153725 4192882 4146641 4207342 4200649 4206232 4192881 4207336 4207337	08-08-78 16-06-81 02-11-79 01-10-80 14-11-77 27-03-79 14-10-80 08-05-79 11-03-80 27-03-79 10-06-80 29-04-80 03-06-80 11-03-80 10-06-80
EP-A- (0274453	13-07-88	FR-A- JP-A-	2609289 63258449	08-07-88 25-10-88
EP-A- ()214639	18-03-87	US-A- US-A- AU-B- AU-A- JP-A-	4599361 4743587 588362 6240886 62103052	08-07-86 10-05-88 14-09-89 12-03-87 13-05-87